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(54) Title: POLYNUCLEOTIDE ENCODING A POLYPEPTIDE HAVING HEPARANASE ACTIVITY AND EXPRESSION OF SAME IN GENETICALLY MODIFIED CELLS

(57) Abstract

A polynucleotide (hpa) encoding a polypeptide having heparanase activity, vectors including same, genetically modified cells expressing heparanase, a recombinant protein having heparanase activity and antisense oligonucleotides and constructs for modulating heparanase expression.

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POLYNUCLEOTIDE ENCODING A POLYPEPTIDE HAVING HEPARANASE ACTIVITY AND EXPRESSION OF SAME IN GENETICALLY MODIFIED CELLS

5 FIELD AND BACKGROUND OF THE INVENTION

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The present invention relates to a polynucleotide, referred to hereinbelow as hpa, encoding a polypeptide having heparanase activity, vectors (nucleic acid constructs) including same and genetically modified cells expressing heparanase. The invention further relates to a recombinant protein having heparanase activity and to antisense oligonucleotides, constructs and ribozymes for down regulating heparanase activity. In addition, the invention relates to heparanase promoter sequences and their uses.

Heparan sulfate proteoglycans: Heparan sulfate proteoglycans (HSPG) are ubiquitous macromolecules associated with the cell surface and extra cellular matrix (ECM) of a wide range of cells of vertebrate and invertebrate tissues (1-4). The basic HSPG structure includes a protein core to which several linear heparan sulfate chains are covalently attached. These polysaccharide chains are typically composed of repeating hexuronic and D-glucosamine disaccharide units that are substituted to a varying extent with N- and O-linked sulfate moieties and N-linked acetyl groups (1-4). Studies on the involvement of ECM molecules in cell attachment, growth and differentiation revealed a central role of HSPG in embryonic morphogenesis, angiogenesis, neurite outgrowth and tissue repair (1-5). HSPG are prominent components of blood vessels (3). In large blood vessels they are concentrated mostly in the intima and inner media, whereas in capillaries they are found mainly in the subendothelial basement membrane where they support proliferating and migrating endothelial cells and stabilize the structure of the capillary wall. The ability of HSPG to interact with ECM macromolecules such as collagen, laminin and fibronectin, and with different attachment sites on plasma membranes suggests a key role for this proteoglycan in the self-assembly and insolubility of ECM components, as well as in cell adhesion and locomotion. Cleavage of the heparan sulfate (HS) chains may therefore result in degradation of the subendothelial ECM and hence may play a decisive role in extravasation of blood-borne cells. HS catabolism is observed in inflammation, wound repair, diabetes, and cancer metastasis, suggesting that enzymes which degrade HS play important roles in pathologic processes. Heparanase activity has been described in activated

immune system cells and highly metastatic cancer cells (6-8), but research has been handicapped by the lack of biologic tools to explore potential causative roles of heparanase in disease conditions.

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Involvement of Heparanase in Tumor Cell Invasion and Metastasis: Circulating tumor cells arrested in the capillary beds of different organs must invade the endothelial cell lining and degrade its underlying basement membrane (BM) in order to invade into the extravascular tissue(s) where they establish metastasis (9, 10). Metastatic tumor cells often attach at or near the intercellular junctions between adjacent endothelial cells. Such attachment of the metastatic cells is followed by rupture of the junctions, retraction of the endothelial cell borders and migration through the breach in the endothelium toward the exposed underlying BM (9). Once located between endothelial cells and the BM, the invading cells must degrade the subendothelial glycoproteins and proteoglycans of the BM in order to migrate out of the vascular compartment. Several cellular enzymes (e.g., collagenase IV, plasminogen activator, cathepsin B, elastase, etc.) are thought to be involved in degradation of BM (10). Among these enzymes is an endo-\beta-Dglucuronidase (heparanase) that cleaves HS at specific intrachain sites (6, 8, 11). Expression of a HS degrading heparanase was found to correlate with the metastatic potential of mouse lymphoma (11), fibrosarcoma and melanoma (8) cells. Moreover, elevated levels of heparanase were detected in sera from metastatic tumor bearing animals and melanoma patients (8) and in tumor biopsies of cancer patients (12).

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The control of cell proliferation and tumor progression by the local microenvironment, focusing on the interaction of cells with the extracellular matrix (ECM) produced by cultured corneal and vascular endothelial cells, was investigated previously by the present inventors. This cultured ECM closely resembles the subendothelium in vivo morphological appearance and molecular composition. It contains collagens (mostly type III and IV, with smaller amounts of types I and V), proteoglycans (mostly heparan sulfate- and dermatan sulfate- proteoglycans, with smaller amounts of chondroitin sulfate proteoglycans), laminin, fibronectin, entactin and elastin (13, 14). The ability of cells to degrade HS in the cultured ECM was studied by allowing cells to interact with a metabolically sulfate labeled ECM, followed by gel filtration (Sepharose 6B) analysis of degradation products released into the culture medium (11). While intact HSPG are eluted next to the void volume of the column

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(Kav<0.2, Mr \sim 0.5x10⁶), labeled degradation fragments of HS side chains are eluted more toward the V_t of the column (0.5<kav<0.8, Mr =5-7x10³) (11).

The heparanase inhibitory effect of various non-anticoagulant species of heparin that might be of potential use in preventing extravasation of blood-borne cells was also investigated by the present inventors. Inhibition of heparanase was best achieved by heparin species containing 16 sugar units or more and having sulfate groups at both the N and O positions. While O-desulfation abolished the heparanase inhibiting effect of heparin, O-sulfated, N-acetylated heparin retained a high inhibitory activity, provided that the N-substituted molecules had a molecular size of about 4,000 daltons or more (7). Treatment of experimental animals with heparanase inhibitors (e.g., non-anticoagulant species of heparin) markedly reduced (>90%) the incidence of lung metastases induced by B16 melanoma, Lewis lung carcinoma and mammary adenocarcinoma cells (7, 8, 16). Heparin fractions with high and low affinity to anti-thrombin III exhibited a comparable high anti-metastatic activity, indicating that the heparanase inhibiting activity of heparin, rather than its anticoagulant activity, plays a role in the anti-metastatic properties of the polysaccharide (7).

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Heparanase activity in the urine of cancer patients: In an attempt to further elucidate the involvement of heparanase in tumor progression and its relevance to human cancer, urine samples for heparanase activity were screened (16a). Heparanase activity was detected in the urine of some, but not all, cancer patients. High levels of heparanase activity were determined in the urine of patients with an aggressive metastatic disease and there was no detectable activity in the urine of healthy donors.

Heparanase activity was also found in the urine of 20% of normal and microalbuminuric insulin dependent diabetes mellitus (IDDM) patients, most likely due to diabetic nephropathy, the most important single disorder leading to renal failure in adults.

Possible involvement of heparanase in tumor angiogenesis: Fibroblast growth factors are a family of structurally related polypeptides characterized by high affinity to heparin (17). They are highly mitogenic for vascular endothelial cells and are among the most potent inducers of neovascularization (17, 18). Basic fibroblast growth factor (bFGF) has been extracted from the subendothelial ECM produced in vitro (19) and from basement membranes of the cornea (20), suggesting that ECM may serve as

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a reservoir for bFGF. Immunohistochemical staining revealed the localization of bFGF in basement membranes of diverse tissues and blood vessels (21). Despite the ubiquitous presence of bFGF in normal tissues, endothelial cell proliferation in these tissues is usually very low, suggesting that bFGF is somehow sequestered from its site of action. Studies on the interaction of bFGF with ECM revealed that bFGF binds to HSPG in the ECM and can be released in an active form by HS degrading enzymes (15, 20, 22). It was demonstrated that heparanase activity expressed by platelets, mast cells, neutrophils, and lymphoma cells is involved in release of active bFGF from ECM and basement membranes (23), suggesting that heparanase activity may not only function in cell migration and invasion, but may also elicit an indirect neovascular response. These results suggest that the ECM HSPG provides a natural storage depot for bFGF and possibly other heparin-binding growth promoting factors (24, 25). Displacement of bFGF from its storage within basement membranes and ECM may therefore provide a novel mechanism for induction of neovascularization in normal

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and pathological situations.

Recent studies indicate that heparin and HS are involved in binding of bFGF to high affinity cell surface receptors and in bFGF cell signaling (26, 27). Moreover, the size of HS required for optimal effect was similar to that of HS fragments released by heparanase (28). Similar results were obtained with vascular endothelial cells growth factor (VEGF) (29), suggesting the operation of a dual receptor mechanism involving HS in cell interaction with heparin-binding growth factors. It is therefore proposed that restriction of endothelial cell growth factors in ECM prevents their systemic action on the vascular endothelium, thus maintaining a very low rate of endothelial cells turnover and vessel growth. On the other hand, release of bFGF from storage in ECM as a complex with HS fragment, may elicit localized endothelial cell proliferation and neovascularization in processes such as wound healing, inflammation and tumor development (24, 25).

Expression of heparanase by cells of the immune system: Heparanase activity correlates with the ability of activated cells of the immune system to leave the circulation and elicit both inflammatory and autoimmune responses. Interaction of platelets, granulocytes, T and B lymphocytes, macrophages and mast cells with the subendothelial ECM is associated with degradation of HS by a specific heparanase activity (6). The enzyme is released from intracellular compartments (e.g., lysosomes,

specific granules, etc.) in response to various activation signals (e.g., thrombin, calcium ionophore, immune complexes, antigens, mitogens, etc.), suggesting its regulated involvement in inflammation and cellular immunity.

Some of the observations regarding the heparanase enzyme were reviewed in reference No. 6 and are listed hereinbelow:

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First, a proteolytic activity (plasminogen activator) and heparanase participate synergistically in sequential degradation of the ECM HSPG by inflammatory leukocytes and malignant cells.

Second, a large proportion of the platelet heparanase exists in a latent form, probably as a complex with chondroitin sulfate. The latent enzyme is activated by tumor cell-derived factor(s) and may then facilitate cell invasion through the vascular endothelium in the process of tumor metastasis.

Third, release of the platelet heparanase from α -granules is induced by a strong stimulant (i.e., thrombin), but not in response to platelet activation on ECM.

Fourth, the neutrophil heparanase is preferentially and readily released in response to a threshold activation and upon incubation of the cells on ECM.

Fifth, contact of neutrophils with ECM inhibited release of noxious enzymes (proteases, lysozyme) and oxygen radicals, but not of enzymes (heparanase, gelatinase) which may enable diapedesis. This protective role of the subendothelial ECM was observed when the cells were stimulated with soluble factors but not with phagocytosable stimulants.

Sixth, intracellular heparanase is secreted within minutes after exposure of T cell lines to specific antigens.

Seventh, mitogens (Con A, LPS) induce synthesis and secretion of heparanase by normal T and B lymphocytes maintained *in vitro*. T lymphocyte heparanase is also induced by immunization with antigen *in vivo*.

Eighth, heparanase activity is expressed by pre-B lymphomas and B-lymphomas, but not by plasmacytomas and resting normal B lymphocytes.

Ninth, heparanase activity is expressed by activated macrophages during incubation with ECM, but there was little or no release of the enzyme into the incubation medium. Similar results were obtained with human myeloid leukemia cells induced to differentiate to mature macrophages.

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Tenth, T-cell mediated delayed type hypersensitivity and experimental autoimmunity are suppressed by low doses of heparanase inhibiting non-anticoagulant species of heparin (30).

Eleventh, heparanase activity expressed by platelets, neutrophils and metastatic tumor cells releases active bFGF from ECM and basement membranes. Release of bFGF from storage in ECM may elicit a localized neovascular response in processes such as wound healing, inflammation and tumor development.

Twelfth, among the breakdown products of the ECM generated by heparanase is a tri-sulfated disaccharide that can inhibit T-cell mediated inflammation in vivo (31). This inhibition was associated with an inhibitory effect of the disaccharide on the production of biologically active TNF α by activated T cells in vitro (31).

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Other potential therapeutic applications: Apart from its involvement in tumor cell metastasis, inflammation and autoimmunity, mammalian heparanase may be applied to modulate: bioavailability of heparin-binding growth factors (15); cellular responses to heparin-binding growth factors (e.g., bFGF, VEGF) and cytokines (IL-8) (31a, 29); cell interaction with plasma lipoproteins (32); cellular susceptibility to certain viral and some bacterial and protozoa infections (33, 33a, 33b); and disintegration of amyloid plaques (34). Heparanase may thus prove useful for conditions such as wound healing, angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases Mammalian heparanase can be used to neutralize plasma heparin, as a potential replacement of protamine. Anti-heparanase antibodies may be applied for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Common use in basic research is expected.

The identification of the *hpa* gene encoding for heparanase enzyme will enable the production of a recombinant enzyme in heterologous expression systems. Availability of the recombinant protein will pave the way for solving the protein structure function relationship and will provide a tool for developing new inhibitors.

Viral Infection: The presence of heparan sulfate on cell surfaces have been shown to be the principal requirement for the binding of Herpes Simplex (33) and Dengue (33a) viruses to cells and for subsequent infection of the cells. Removal of the cell surface heparan sulfate by heparanase may

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therefore abolish virus infection. In fact, treatment of cells with bacterial heparitinase (degrading heparan sulfate) or heparinase (degrading heparan) reduced the binding of two related animal herpes viruses to cells and rendered the cells at least partially resistant to virus infection (33). There are some indications that the cell surface heparan sulfate is also involved in HIV infection (33b).

Neurodegenerative diseases: Heparan sulfate proteoglycans were identified in the prion protein amyloid plaques of Genstmann-Straussler Syndrome, Creutzfeldt-Jakob disease and Scrape (34). Heparanase may disintegrate these amyloid plaques which are also thought to play a role in the pathogenesis of Alzheimer's disease.

Restenosis and Atherosclerosis: Proliferation of arterial smooth muscle cells (SMCs) in response to endothelial injury and accumulation of cholesterol rich lipoproteins are basic events in the pathogenesis of atherosclerosis and restenosis (35). Apart from its involvement in SMC proliferation (i.e., low affinity receptors for heparin-binding growth factors), HS is also involved in lipoprotein binding, retention and uptake (36). It was demonstrated that HSPG and lipoprotein lipase participate in a novel catabolic pathway that may allow substantial cellular and interstitial accumulation of cholesterol rich lipoproteins (32). The latter pathway is expected to be highly atherogenic by promoting accumulation of apoB and apoE rich lipoproteins (i.e. LDL, VLDL, chylomicrons), independent of feed back inhibition by the cellular sterol content. Removal of SMC HS by heparanase is therefore expected to inhibit both SMC proliferation and lipid accumulation and thus may halt the progression of restenosis and atherosclerosis.

Gene therapy:

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The ultimate goal in the management of inherited as well as acquired diseases is a rational therapy with the aim to eliminate the underlying biochemical defects associated with the disease rather then symptomatic treatment. Gene therapy is a promising candidate to meet these objectives. Initially it was developed for treatment of genetic disorders, however, the consensus view today is that it offers the prospect of providing therapy for a variety of acquired diseases, including cancer, viral infections, vascular diseases and neurodegenerative disorders.

The gene-based therapeutic can act either intracellularly, affecting only the cells to which it is delivered, or extracellularly, using the recipient cells as local endogenous factories for the therapeutic product(s). The

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application of gene therapy may follow any of the following strategies: (i) prophylactic gene therapy, such as using gene transfer to protect cells against viral infection; (ii) cytotoxic gene therapy, such as cancer therapy, where genes encode cytotoxic products to render the target cells vulnerable to attack by the normal immune response; (iii) biochemical correction, primarily for the treatment of single gene defects, where a normal copy of the gene is added to the affected or other cells.

To allow efficient transfer of the therapeutic genes, a variety of gene delivery techniques have been developed based on viral and non-viral vector systems. The most widely used and most efficient systems for delivering genetic material into target cells are viral vectors. So far, 329 clinical studies (phase I, I/II and II) with over 2,500 patients have been initiated Worldwide since 1989 (50).

The approach of gene addition pose serious barriers. The expression of many genes is tightly regulated and context dependent, so achieving the correct balance and function of expression is challenging. The gene itself is often quite large, containing many exons and introns. The delivery vector is usually a virus, which can infect with a high efficiency but may, on the other hand, induce immunological response and consequently decreases effectiveness, especially upon secondary administration. Most of the current expression vector-based gene therapy protocols fail to achieve clinically significant transgene expression required for treating genetic diseases. Apparently, it is difficult to deliver enough virus to the right cell type to elicit an effective and therapeutic effect (51)

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Homologous recombination, which was initially considered to be of limited use for gene therapy because of its low frequency in mammalian cells, has recently emerged as a potential strategy for developing gene therapy. Different approaches have been used to study homologous recombination in mammalian cells; some involve DNA repair mechanisms. These studies aimed at either gene disruption or gene correction and include RNA/DNA chimeric oligonucleotides, small or large homologous DNA fragments, or adeno-associated viral vectors. Most of these studies show a reasonable frequency of homologous recombination, which warrants further in vivo testing (52). Homologous recombination-based gene therapy has the potential to develop into a powerful therapeutic modality for genetic diseases. It can offer permanent expression and normal regulation of corrected genes in appropriate cells or organs and probably can be used for treating dominantly inherited diseases such as polycystic kidney disease.

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Genomic sequences function in regulation of gene expression:

The efficient expression of therapeutic genes in target cells or tissues is an important component of efficient and safe gene therapy. The expression of genes is driven by the promoter region upstream of the coding sequence, although regulation of expression may be supplemented by farther upstream or downstream DNA sequences or DNA in the introns of the gene. Since this important information is embedded in the DNA, the description of gene structure is crucial to the analysis of gene regulation. Characterization of cell specific or tissue specific promoters, as well as other tissue specific regulatory elements enables the use of such sequences to direct efficient cell specific, or developmental stage specific gene expression. This information provides the basis for targeting individual genes and for control of their expression by exogenous agents, such as drugs. Identification of transcription factors and other regulatory proteins required for proper gene expression will point at new potential targets for modulating gene expression, when so desired or required.

Efficient expression of many mammalian genes depends on the presence of at least one intron. The expression of mouse thymidylate synthase (TS) gene, for example, is greatly influenced by intron sequences. The addition of almost any of the introns from the mouse TS gene to an intronless TS minigene leads to a large increase in expression (42). The involvement of intron 1 in the regulation of expression was demonstrated for many other genes. In human factor IX (hFIX), intron 1 is able to increase the expression level about 3 fold mare as compared to that of the hFIX cDNA (43). The expression enhancing activity of intron 1 is due to efficient functional splicing sequences, present in the precursor mRNA. By being efficiently assembled into spliceosome complexes, transcripts with splicing sequences may be better protected in the nucleus from random degradations, than those without such sequences (44).

A forward-inserted intron1-carrying hFIX expression cassette suggested to be useful for directed gene transfer, while for retroviral-mediated gene transfer system, reversely-inserted intron 1-carrying hFIX expression cassette was considered (43).

A highly conserved cis-acting sequence element was identified in the first intron of the mouse and rat c-Ha-ras, and in the first exon of Ha- and Ki-ras genes of human, mouse and rat. This cis-acting regulatory sequence confers strong transcription enhancer activity that is differentially modulated by steroid hormones in metastatic and nonmetastatic

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subpopulations. Perturbations in the regulatory activities of such cis-acting sequences may play an important role in governing oncogenic potency of Ha-ras through transcriptional control mechanisms (45).

Intron sequences affect tissue specific, as well as inducible gene expression. A 182 bp intron 1 DNA segment of the mouse Col2a1 gene contains the necessary information to confer high-level, temporally correct, chondrocyte expression on a reporter gene in intact mouse embryos, while Col2a1 promoter sequences are dispensable for chondrocyte expression (46). In Col1A1 gene the intron plays little or no role in constitutive expression of collagen in the skin, and in cultured cells derived from the skin, however, in the lungs of young mice, intron deletion results in decrease of expression to less than 50 % (47).

A classical enhancer activity was shown in the 2 kb intron fragment in bovine beta-casein gene. The enhancer activity was largely dependent on the lactogenic hormones, especially prolactin. It was suggested that several elements in the intron-1 of the bovine beta-casein gene cooperatively interact not only with each other but also with its promoter for hormonal induction (48).

Identification and characterization of regulatory elements in genomic non-coding sequences, such as introns, provides a tool for designing and constructing novel vectors for tissue specific, hormone regulated or any other defined expression pattern, for gene therapy. Such an expression cassette was developed, utilizing regulatory elements from the human cytokeratin 18 (K18) gene, including 5' genomic sequences and one of its introns. This cassette efficiently expresses reporter genes, as well as the human cystic fibrosis transmembrane conductance regulator (CFTR) gene, in cultured lung epithelial cells (49).

Alternative splicing:

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Alternative splicing of pre mRNA is a powerful and versatile regulatory mechanism that can effect quantitative control of gene expression and functional diversification of proteins. It contributes to major developmental decisions and also to a fine-tuning of gene function. Genetic and biochemical approaches have identified cis-acting regulatory elements and trans-acting factors that control alternative splicing of specific mRNAs. This mechanism results in the generation of variant isoforms of various proteins from a single gene. These include cell surface molecules such as CD44, receptors, cytokines such as VEGF and enzymes. Products of

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alternatively spliced transcripts differ in their expression pattern, substrate specificity and other biological parameters.

The FGF receptor RNA undergoes alternative splicing which results in the production of several isoforms, which exhibit different ligand binding specificities. The alternative splicing is regulated in a cell specific manner (53).

Alternative spliced mRNAs are often correlated with malignancy. An increase in specific splice variant of tyrosinase was identified in murine melanomas (54). Multiple splicing variants of estrogen receptor are present in individual human breast tumors. CD44 has various isoform, some are characteristic of malignant tissues.

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Identification of tumor specific alternative splice variants provide new tool for cancer diagnostics. CD44 variants have been used for detection of malignancy in urine samples from patients with urothelial cancer by competitive RT-PCR (55). CD44 exon 6 was suggested as prognostic indicator of metastasis in breast cancer (56).

Different enzymes or polypeptides generated by alternative splicing may have different function or catalytic specificity. The identification and characterization of the enzyme forms, which are involved in pathological processes, is crucial for the design of appropriate and efficient drugs.

Modulation of gene expression - Antisense technology:

An antisense oligonucleotide (e.g., antisense oligodeoxyribonucleotide) may bind its target nucleic acid either by Watson-Crick base pairing or Hoogsteen and anti-Hoogsteen base pairing (64). According to the Watson-Crick base pairing, heterocyclic bases of the antisense oligonucleotide form hydrogen bonds with the heterocyclic bases of target single-stranded nucleic acids (RNA or single-stranded DNA), whereas according to the Hoogsteen base pairing, the heterocyclic bases of the target nucleic acid are double-stranded DNA, wherein a third strand is accommodated in the major groove of the B-form DNA duplex by Hoogsteen and anti-Hoogsteen base pairing to form a triple helix structure.

According to both the Watson-Crick and the Hoogsteen base pairing models, antisense oligonucleotides have the potential to regulate gene expression and to disrupt the essential functions of the nucleic acids in cells. Therefore, antisense oligonucleotides have possible uses in modulating a wide range of diseases in which gene expression is altered.

Since the development of effective methods for chemically synthesizing oligonucleotides, these molecules have been extensively used

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in biochemistry and biological research and have the potential use in medicine, since carefully devised oligonucleotides can be used to control gene expression by regulating levels of transcription, transcripts and/or translation.

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Oligodeoxyribonucleotides as long as 100 base pairs (bp) are routinely synthesized by solid phase methods using commercially available, fully automated synthesis machines. The chemical synthesis of oligoribonucleotides, however, is far less routine. Oligoribonucleotides are also much less stable than oligodeoxyribonucleotides, a fact which has contributed to the more prevalent use of oligodeoxyribonucleotides in medical and biological research, directed at, for example, the regulation of transcription or translation levels.

Gene expression involves few distinct and well regulated steps. The first major step of gene expression involves transcription of a messenger RNA (mRNA) which is an RNA sequence complementary to the antisense (i.e., -) DNA strand, or, in other words, identical in sequence to the DNA sense (i.e., +) strand, composing the gene. In eukaryotes, transcription occurs in the cell nucleus.

The second major step of gene expression involves translation of a protein (e.g., enzymes, structural proteins, secreted proteins, gene expression factors, etc.) in which the mRNA interacts with ribosomal RNA complexes (ribosomes) and amino acid activated transfer RNAs (tRNAs) to direct the synthesis of the protein coded for by the mRNA sequence.

Initiation of transcription requires specific recognition of a promoter DNA sequence located upstream to the coding sequence of a gene by an RNA-synthesizing enzyme -- RNA polymerase. This recognition is preceded by sequence-specific binding of one or more transcription factors to the promoter sequence. Additional proteins which bind at or close to the promoter sequence may trans upregulate transcription via cis elements known as enhancer sequences. Other proteins which bind to or close to the promoter, but whose binding prohibits the action of RNA polymerase, are known as repressors.

There are also evidence that in some cases gene expression is downregulated by endogenous antisense RNA repressors that bind a complementary mRNA transcript and thereby prevent its translation into a functional protein.

Thus, gene expression is typically upregulated by transcription factors and enhancers and downregulated by repressors.

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However, in many disease situation gene expression is impaired. In many cases, such as different types of cancer, for various reasons the expression of a specific endogenous or exogenous (e.g., of a pathogen such as a virus) gene is upregulated. Furthermore, in infectious diseases caused by pathogens such as parasites, bacteria or viruses, the disease progression depends on expression of the pathogen genes, this phenomenon may also be considered as far as the patient is concerned as upregulation of exogenous genes.

Most conventional drugs function by interaction with and modulation of one or more targeted endogenous or exogenous proteins, e.g., enzymes. Such drugs, however, typically are not specific for targeted proteins but interact with other proteins as well. Thus, a relatively large dose of drug must be used to effectively modulate a targeted protein.

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Typical daily doses of drugs are from 10-5 - 10-1 millimoles per kilogram of body weight or 10-3 - 10 millimoles for a 100 kilogram person. If this modulation instead could be effected by interaction with and inactivation of mRNA, a dramatic reduction in the necessary amount of drug could likely be achieved, along with a corresponding reduction in side effects. Further reductions could be effected if such interaction could be rendered site-specific. Given that a functioning gene continually produces mRNA, it would thus be even more advantageous if gene transcription could be arrested in its entirety.

Given these facts, it would be advantageous if gene expression could be arrested or downmodulated at the transcription level.

The ability of chemically synthesizing oligonucleotides and analogs thereof having a selected predetermined sequence offers means for downmodulating gene expression. Three types of gene expression modulation strategies may be considered.

At the transcription level, antisense or sense oligonucleotides or analogs that bind to the genomic DNA by strand displacement or the formation of a triple helix, may prevent transcription (64).

At the transcript level, antisense oligonucleotides or analogs that bind target mRNA molecules lead to the enzymatic cleavage of the hybrid by intracellular RNase H (65). In this case, by hybridizing to the targeted mRNA, the oligonucleotides or oligonucleotide analogs provide a duplex hybrid recognized and destroyed by the RNase H enzyme. Alternatively, such hybrid formation may lead to interference with correct splicing (66).

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As a result, in both cases, the number of the target mRNA intact transcripts ready for translation is reduced or eliminated.

At the translation level, antisense oligonucleotides or analogs that bind target mRNA molecules prevent, by steric hindrance, binding of essential translation factors (ribosomes), to the target mRNA, a phenomenon known in the art as hybridization arrest, disabling the translation of such mRNAs (67).

Thus, antisense sequences, which as described hereinabove may arrest the expression of any endogenous and/or exogenous gene depending on their specific sequence, attracted much attention by scientists and pharmacologists who were devoted at developing the antisense approach into a new pharmacological tool (68).

For example, several antisense oligonucleotides have been shown to arrest hematopoietic cell proliferation (69), growth (70), entry into the S phase of the cell cycle (71), reduced survival (72) and prevent receptor mediated responses (73). For use of antisense oligonucleotides as antiviral agents the reader is referred to reference 74.

For efficient *in vivo* inhibition of gene expression using antisense oligonucleotides or analogs, the oligonucleotides or analogs must fulfill the following requirements (i) sufficient specificity in binding to the target sequence; (ii) solubility in water; (iii) stability against intra- and extracellular nucleases; (iv) capability of penetration through the cell membrane; and (v) when used to treat an organism, low toxicity.

Unmodified oligonucleotides are impractical for use as antisense sequences since they have short *in vivo* half-lives, during which they are degraded rapidly by nucleases. Furthermore, they are difficult to prepare in more than milligram quantities. In addition, such oligonucleotides are poor cell membrane penetraters (75).

Thus it is apparent that in order to meet all the above listed requirements, oligonucleotide analogs need to be devised in a suitable manner. Therefore, an extensive search for modified oligonucleotides has been initiated.

For example, problems arising in connection with double-stranded DNA (dsDNA) recognition through triple helix formation have been diminished by a clever "switch back" chemical linking, whereby a sequence of polypurine on one strand is recognized, and by "switching back", a homopurine sequence on the other strand can be recognized. Also, good

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helix formation has been obtained by using artificial bases, thereby improving binding conditions with regard to ionic strength and pH.

In addition, in order to improve half-life as well as membrane penetration, a large number of variations in polynucleotide backbones have been done, nevertheless with little success.

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Oligonucleotides can be modified either in the base, the sugar or the phosphate moiety. These modifications include, for example, the use of methylphosphonates, monothiophosphates, dithiophosphates, phosphoramidates, phosphate esters, bridged phosphorothioates, bridged phosphoramidates, bridged methylenephosphonates, dephospho internucleotide analogs with siloxane bridges, carbonate bridges, carboxymethyl ester bridges, carboxymethyl ester bridges, carboxymethyl ester bridges, acetamide bridges, carboxamate bridges, thioether bridges, sulfoxy bridges, sulfono bridges, various "plastic" DNAs, \alpha-anomeric bridges and borane derivatives. For further details the reader is referred to reference 76.

International patent application WO 89/12060 discloses various building blocks for synthesizing oligonucleotide analogs, as well as oligonucleotide analogs formed by joining such building blocks in a defined sequence. The building blocks may be either "rigid" (i.e., containing a ring structure) or "flexible" (i.e., lacking a ring structure). In both cases, the building blocks contain a hydroxy group and a mercapto group, through which the building blocks are said to join to form oligonucleotide analogs. The linking moiety in the oligonucleotide analogs is selected from the group consisting of sulfide (-S-), sulfoxide (-SO-), and sulfone (-SO₂-). However, the application provides no data supporting the specific binding of an oligonucleotide analog to a target oligonucleotide.

International patent application WO 92/20702 describe an acyclic oligonucleotide which includes a peptide backbone on which any selected chemical nucleobases or analogs are stringed and serve as coding characters as they do in natural DNA or RNA. These new compounds, known as peptide nucleic acids (PNAs), are not only more stable in cells than their natural counterparts, but also bind natural DNA and RNA 50 to 100 times more tightly than the natural nucleic acids cling to each other (77). PNA oligomers can be synthesized from the four protected monomers containing thymine, cytosine, adenine and guanine by Merrifield solid-phase peptide synthesis. In order to increase solubility in water and to prevent aggregation, a lysine amide group is placed at the C-terminal.

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Thus, antisense technology requires pairing of messenger RNA with an oligonucleotide to form a double helix that inhibits translation. The concept of antisense-mediated gene therapy was already introduced in 1978 for cancer therapy. This approach was based on certain genes that are crucial in cell division and growth of cancer cells. Synthetic fragments of genetic substance DNA can achieve this goal. Such molecules bind to the targeted gene molecules in RNA of tumor cells, thereby inhibiting the translation of the genes and resulting in dysfunctional growth of these cells. Other mechanisms has also been proposed. These strategies have been used, with some success in treatment of cancers, as well as other illnesses, including viral and other infectious diseases. Antisense oligonucleotides are typically synthesized in lengths of 13-30 nucleotides. The life span of oligonucleotide molecules in blood is rather short. Thus, they have to be chemically modified to prevent destruction by ubiquitous nucleases present in the body. Phosphorothioates are very widely used modification in antisense oligonucleotide ongoing clinical trials (57). A new generation of antisense molecules consist of hybrid antisense oligonucleotide with a central portion of synthetic DNA while four bases on each end have been modified with 2'O-methyl ribose to resemble RNA. In preclinical studies in laboratory animals, such compounds have demonstrated greater stability to metabolism in body tissues and an improved safety profile when compared with the first-generation unmodified phosphorothioate (Hybridon Inc. Dosens of other nucleotide analogs have also been tested in news). antisense technology.

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RNA oligonucleotides may also be used for antisense inhibition as they form a stable RNA-RNA duplex with the target, suggesting efficient inhibition. However, due to their low stability RNA oligonucleotides are typically expressed inside the cells using vectors designed for this purpose. This approach is favored when attempting to target a mRNA that encodes an abundant and long-lived protein (57).

Recent scientific publications have validated the efficacy of antisense compounds in animal models of hepatitis, cancers, coronary artery restenosis and other diseases. The first antisense drug was recently approved by the FDA. This drug Fomivirsen, developed by Isis, is indicated for local treatment of cytomegalovirus in patients with AIDS who are intolerant of or have a contraindication to other treatments for CMV retinitis or who were insufficiently responsive to previous treatments for CMV retinitis (Pharmacotherapy News Network).

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Several antisense compounds are now in clinical trials in the United States. These include locally administered antivirals, systemic cancer therapeutics. Antisense therapeutics has the potential to treat many life-threatening diseases with a number of advantages over traditional drugs. Traditional drugs intervene after a disease-causing protein is formed. Antisense therapeutics, however, block mRNA transcription/translation and intervene before a protein is formed, and since antisense therapeutics target only one specific mRNA, they should be more effective with fewer side effects than current protein-inhibiting therapy.

A second option for disrupting gene expression at the level of transcription uses synthetic oligonucleotides capable of hybridizing with double stranded DNA. A triple helix is formed. Such oligonucleotides may prevent binding of transcription factors to the gene's promoter and therefore inhibit transcription. Alternatively, they may prevent duplex unwinding and, therefore, transcription of genes within the triple helical structure.

Another approach is the use of specific nucleic acid sequences to act as decoys for transcription factors. Since transcription factors bind specific DNA sequences it is possible to synthesize oligonucleotides that will effectively compete with the native DNA sequences for available transcription factors *in vivo*. This approach requires the identification of gene specific transcription factor (57).

Indirect inhibition of gene expression was demonstrated for matrix metalloproteinase genes (MMP-1, -3, and -9), which are associated with invasive potential of human cancer cells. E1AF is a transcription activator of MMP genes. Expression of E1AF antisense RNA in HSC3AS cells showed decrease in mRNA and protein levels of MMP-1, -3, and -9. Moreover, HSC3AS showed lower invasive potential in vitro and *in vivo*. These results imply that transfection of antisense inhibits tumor invasion by down-regulating MMP genes (58).

Ribozymes:

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Ribozymes are being increasingly used for the sequence-specific inhibition of gene expression by the cleavage of mRNAs encoding proteins of interest. The possibility of designing ribozymes to cleave any specific target RNA has rendered them valuable tools in both basic research and therapeutic applications. In the therapeutics area, ribozymes have been exploited to target viral RNAs in infectious diseases, dominant oncogenes in cancers and specific somatic mutations in genetic disorders. Most notably, several ribozyme gene therapy protocols for HIV patients are

already in Phase 1 trials (62). More recently, ribozymes have been used for transgenic animal research, gene target validation and pathway elucidation. Several ribozymes are in various stages of clinical trials. ANGIOZYME was the first chemically synthesized ribozyme to be studied in human clinical trials. ANGIOZYME specifically inhibits formation of the VEGF-r (Vascular Endothelial Growth Factor receptor), a key component in the angiogenesis pathway. Ribozyme Pharmaceuticals, Inc., as well as other firms have demonstrated the importance of anti-angiogenesis therapeutics in animal models. HEPTAZYME, a ribozyme designed to selectively destroy Hepatitis C Virus (HCV) RNA, was found effective in decreasing Hepatitis C viral RNA in cell culture assays (Ribozyme Pharmaceuticals, Incorporated - WEB home page).

Gene disruption in animal models:

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The emergence of gene inactivation by homologous recombination methodology in embryonic stem cells has revolutionized the field of mouse genetics. The availability of a rapidly growing number of mouse null mutants has represented an invaluable source of knowledge on mammalian development, cellular biology and physiology, and has provided many models for human inherited diseases. Animal models are required for an effective drug delivery development program and evaluation of gene therapy approach. The improvement of the original knockout strategy, as well as exploitation of exogenous enzymatic systems that are active in the recombination process, has been considerably extended the range of genetic manipulations that can be produced. Additional methods have been developed to provide versatile research tools: Double replacement method, sequential gene targeting, conditional cell type specific gene targeting, single copy integration method, inducible gene targeting, gene disruption by viral delivery, replacing one gene with another, the so called knock-in method and the induction of specific balanced chromosomal translocation. It is now possible to introduce a point mutation as a unique change in the entire genome, therefore allowing very fine dissection of gene function in Furthermore, the advent of methods allowing conditional gene targeting opens the way for analysis of consequence of a particular mutation in a defined organ and at a specific time during the life of the experimental animal (59).

DNA vaccination:

Observations in the early 1990s that plasmid DNA could directly transfect animal cells in vivo sparked exploration of the use of DNA

plasmids to induce immune response by direct injection into animal of DNA encoding antigenic protein. When a DNA vaccine plasmid enters the eukaryotic cell, the protein it encodes is transcribed and translated within the cell. In the case of pathogens, these proteins are presented to the immune system in their native form, mimicking the presentation of antigens during a natural infection. DNA vaccination is particularly useful for the induction of T cell activation. It was applied for viral and bacterial infectious diseases, as well as for allergy and for cancer. The central hypothesis behind active specific immunotherapy for cancer is that tumor cells express unique antigens that should stimulate the immune system. The first DNA vaccine against tumor was carcino-embrionic antigen (CEA). DNA vaccinated animals expressed immunoprotection and immunotherapy of human CEA-expressing syngeneic mouse colon and breast carcinoma (61). In a mouse model of neuroblastoma, DNA immunization with HuD resulted in tumor growth inhibition with no neurological disease (60). Immunity to the brown locus protein, gp⁷⁵ tyrosinase-related protein-1, associated with melanoma, was investigated in a syngeneic mouse model. Priming with human gp75 DNA broke tolerance to mouse gp75. Immunity against mouse gp75 provided significant tumor protection (60).

Glycosyl hydrolases:

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Glycosyl hydrolases are a widespread group of enzymes that hydrolyze the o-glycosidic bond between two or more carbohydrates or between a carbohydrate and a noncarbohydrate moiety. The enzymatic hydrolysis of glycosidic bond occurs by using major one or two mechanisms leading to overall retention or inversion of the anomeric configuration. In both mechanisms catalysis involves two residues: a proton donor and a nucleophile. Glycosyl hydrolyses have been classified into 58 families based on amino acid similarities. The glycosyl hydrolyses from families 1, 2, 5, 10, 17, 30, 35, 39 and 42 act on a large variety of substrates, however, they all hydrolyze the glycosidic bond in a general acid catalysis mechanism, with retention of the anomeric configuration. The mechanism involves two glutamic acid residues, which are the proton donors and the nucleophile, with an aspargine always preceding the proton donor. Analyses of a set of known 3D structures from this group revealed that their catalytic domains, despite the low level of sequence identity, adopt a similar (α/β) 8 fold with the proton donor and the nucleophile located at the C-terminal ends of strands $\beta4$ and $\beta7$, respectively. Mutations

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in the functional conserved amino acids of lysosomal glycosyl hydrolases were identified in lysosomal storage diseases.

Lysosomal glycosyl hydrolases including β -glucuronidase, β -manosidase, β -glucocerebrosidase, β -galactosidase and α -L iduronidase, are all exo-glycosyl hydrolases, belong to the GH-A clan and share a similar catalytic site. However, many endo-glucanases from various organisms, such as bacterial and fungal xylenases and cellulases share this catalytic domain.

Genomic sequence of hpa gene and its implications:

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It is well established that heparanase activity is correlated with This correlation was demonstrated at the level of cancer metastasis. enzymatic activity as well as the levels of protein and hpa cDNA expression in highly metastatic cancer cells as compared with non-metastatic cells. As such, inhibition of heparanase activity is desirable, and has been attempted by several means. The genomic region, encoding the hpa gene and the surrounding, provides a new powerful tool for regulation of heparanase activity at the level of gene expression. Regulatory sequences may reside in noncoding regions both upstream and downstream the transcribed region as well as in intron sequences. A DNA sequence upstream of the transcription start site contains the promoter region and potential regulatory elements. Regulatory factors, which interact with the promoter region may be identified and be used as potential drugs for inhibition of cancer, metastasis and inflammation. The promoter region can be used to screen for inhibitors of heparanase gene expression. Furthermore, the hpa promoter can be used to direct cell specific, particularly cancer cell specific, expression of foreign genes, such as cytotoxic or apoptotic genes, in order to specifically destroy cancer cells.

Cancer and yet unknown related genetic disorders may involve rearrangements and mutations in the heparanase gene, either in coding or non-coding regions. Such mutations may affect expression level or enzymatic activity. The genomic sequence of hpa enables the amplification of specific genomic DNA fragments, identification and diagnosis of mutations.

There is thus a widely recognized need for, and it would be highly advantageous to have genomic, cDNA and composite polynucleotides encoding a polypeptide having heparanase activity, vectors including same, genetically modified cells expressing heparanase and a recombinant protein

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having heparanase activity, as well as antisense oligonucleotides, constructs and ribozymes which can be used for down regulation heparanase activity.

SUMMARY OF THE INVENTION

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Cloning of the human hpa gene which encodes heparanase, and expression of recombinant heparanase by transfected host cells is reported herein, as well as downregulation of heparanase activity by antisense technology.

A purified preparation of heparanase isolated from human hepatoma cells was subjected to tryptic digestion and microsequencing. The YGPDVGQPR (SEQ ID NO:8) sequence revealed was used to screen EST databases for homology to the corresponding back translated DNA sequence. Two closely related EST sequences were identified and were thereafter found to be identical. Both clones contained an insert of 1020 bp which included an open reading frame of 973 bp followed by a 27 bp of 3' untranslated region and a Poly A tail. Translation start site was not identified.

Cloning of the missing 5' end of hpa was performed by PCR amplification of DNA from placenta Marathon RACE cDNA composite using primers selected according to the EST clones sequence and the linkers of the composite. A 900 bp PCR fragment, partially overlapping with the identified 3' encoding EST clones was obtained. The joined cDNA fragment (hpa), 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons.

Cloning an extended 5' sequence was enabled from the human SK-hep1 cell line by PCR amplification using the Marathon RACE. The 5' extended sequence of the SK-hep1 hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQ ID NO:9). The assembled sequence contained an open reading frame, SEQ ID NOs: 13 and 15, which encodes, as shown in SEQ ID NOs:14 and 15, a polypeptide of 592 amino acids with a calculated molecular weight of 66,407 daltons.

The ability of the *hpa* gene product to catalyze degradation of heparan sulfate in an *in vitro* assay was examined by expressing the entire open reading frame of *hpa* in insect cells, using the Baculovirus expression system. Extracts and conditioned media of cells infected with virus containing the *hpa* gene, demonstrated a high level of heparan sulfate degradation activity both towards soluble ECM-derived HSPG and intact

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ECM. This degradation activity was inhibited by heparin, which is another substrate of heparanase. Cells infected with a similar construct containing no hpa gene had no such activity, nor did non-infected cells. The ability of heparanase expressed from the extended 5' clone towards heparin was demonstrated in a mammalian expression system.

The expression pattern of hpa RNA in various tissues and cell lines was investigated using RT-PCR. It was found to be expressed only in tissues and cells previously known to have heparanase activity.

A panel of monochromosomal human/CHO and human/mouse somatic cell hybrids was used to localize the human heparanase gene to human chromosome 4. The newly isolated heparanase sequence can be used to identify a chromosome region harboring a human heparanase gene in a chromosome spread.

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A human genomic library was screened and the human locus harboring the heparanase gene isolated, sequenced and characterized. Alternatively spliced heparanase mRNAs were identified and characterized. The human heparanase promoter has been isolated, identified and positively tested for activity. The mouse heparanase promoter has been isolated and identified as well. Antisense heparanase constructs were prepared and their influence on cells *in vitro* tested. A predicted heparanase active site was identified. And finally, the presence of sequences hybridizing with human heparanase sequences was demonstrated for a variety of mammalians and for an avian.

According to one aspect of the present invention there is provided an isolated nucleic acid comprising a genomic, complementary or composite polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

According to further features in preferred embodiments of the invention described below, the polynucleotide or a portion thereof is hybridizable with SEQ ID NOs: 9, 13, 42, 43 or a portion thereof at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 μ g/ml salmon sperm DNA, and ³²p labeled probe and wash at 68 °C with 3 x SSC and 0.1 % SDS.

According to still further features in the described preferred embodiments the polynucleotide or a portion thereof is at least 60 % identical with SEQ ID NOs: 9, 13, 42, 43 or portions thereof as determined using the Bestfit procedure of the DNA sequence analysis software package

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developed by the Genetic Computer Group (GCG) at the university of Wisconsin (gap creation penalty - 12, gap extension penalty - 4).

According to still further features in the described preferred embodiments the polypeptide is as set forth in SEQ ID NOs:10, 14, 44 or portions thereof.

According to still further features in the described preferred embodiments the polypeptide is at least 60 % homologous to SEQ ID NOs:10, 14, 44 or portions thereof as determined with the Smith-Waterman algorithm, using the Bioaccelerator platform developed by Compugene (gapop: 10.0, gapext: 0.5, matrix: blosum62).

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According to additional aspects of the present invention there are provided a nucleic acid construct (vector) comprising the isolated nucleic acid described herein and a host cell comprising the construct.

According to a further aspect of the present invention there is provided an antisense oligonucleotide comprising a polynucleotide or a polynucleotide analog of at least 10 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity.

According to an additional aspect of the present invention there is provided a method of *in vivo* downregulating heparanase activity comprising the step of *in vivo* administering the antisense oligonucleotide herein described.

According to yet an additional aspect of the present invention there is provided a pharmaceutical composition comprising the antisense oligonucleotide herein described and a pharmaceutically acceptable carrier.

According to still an additional aspect of the present invention there is provided a ribozyme comprising the antisense oligonucleotide described herein and a ribozyme sequence.

According to a further aspect of the present invention there is provided an antisense nucleic acid construct comprising a promoter sequence and a polynucleotide sequence directing the synthesis of an antisense RNA sequence of at least 10 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity.

According to further features in preferred embodiments of the invention described below, the polynucleotide strand encoding the polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 9, 13, 42 or 43.

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According to still further features in the described preferred embodiments the polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 10, 14 or 44.

According to still a further aspect of the present invention there is provided a method of *in vivo* downregulating heparanase activity comprising the step of *in vivo* administering the antisense nucleic acid construct herein described.

According to yet a further aspect of the present invention there is provided a pharmaceutical composition comprising the antisense nucleic acid construct herein described and a pharmaceutically acceptable carrier.

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According to a further aspect of the present invention there is provided a nucleic acid construct comprising a polynucleotide sequence functioning as a promoter, the polynucleotide sequence is derived from SEQ ID NO:42 and includes at least nucleotides 2535-2635 thereof or from SEQ ID NO:43 and includes at least nucleotides 320-420.

According to a further aspect of the present invention there is provided a method of expressing a polynucleotide sequence comprising the step of ligating the polynucleotide sequence into the nucleic acid construct described above, downstream of the polynucleotide sequence derived from SEQ ID NOs:42 or 43.

According to a further aspect of the present invention there is provided a recombinant protein comprising a polypeptide having heparanase catalytic activity.

According to further features in preferred embodiments of the invention described below, the polypeptide includes at least a portion of SEQ ID NOs:10, 14 or 44.

According to still further features in the described preferred embodiments the protein is encoded by a polynucleotide hybridizable with SEQ ID NOs: 9, 13, 42, 43 or a portion thereof at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 μ g/ml salmon sperm DNA, and ³²p labeled probe and wash at 68 °C with 3 x SSC and 0.1 % SDS.

According to still further features in the described preferred embodiments the protein is encoded by a polynucleotide at least 60 % identical with SEQ ID NOs: 9, 13, 42, 43 or portions thereof as determined using the Bestfit procedure of the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the university of Wisconsin (gap creation penalty - 12, gap extension penalty - 4).

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According to a further aspect of the present invention there is provided a pharmaceutical composition comprising, as an active ingredient, the recombinant protein herein described.

According to a further aspect of the present invention there is provided a method of identifying a chromosome region harboring a heparanase gene in a chromosome spread comprising the steps of (a) hybridizing the chromosome spread with a tagged polynucleotide probe encoding heparanase; (b) washing the chromosome spread, thereby removing excess of non-hybridized probe; and (c) searching for signals associated with the hybridized tagged polynucleotide probe, wherein detected signals being indicative of a chromosome region harboring a heparanase gene.

According to a further aspect of the present invention there is provided a method of *in vivo* eliciting anti-heparanase antibodies comprising the steps of administering a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*. Accordingly, there is provided also a DNA vaccine for *in vivo* eliciting anti-heparanase antibodies comprising a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*.

The present invention can be used to develop new drugs to inhibit tumor cell metastasis, inflammation and autoimmunity. The identification of the *hpa* gene encoding for heparanase enzyme enables the production of a recombinant enzyme in heterologous expression systems. Additional features, advantages, uses and applications of the present invention in biological science and in diagnostic and therapeutic medicine are described hereinafter.

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BRIEF DESCRIPTION OF THE DRAWINGS

The invention herein described, by way of example only, with reference to the accompanying drawings, wherein:

FIG. 1 presents nucleotide sequence and deduced amino acid sequence of hpa cDNA. A single nucleotide difference at position 799 (A to T) between the EST (Expressed Sequence Tag) and the PCR amplified cDNA (reverse transcribed RNA) and the resulting amino acid substitution (Tyr to Phe) are indicated above and below the substituted unit,

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respectively. Cysteine residues and the poly adenylation consensus sequence are underlined. The asterisk denotes the stop codon TGA.

FIG. 2 demonstrates degradation of soluble sulfate labeled HSPG substrate by lysates of High Five cells infected with pFhpa2 virus. Lysates of High Five cells that were infected with pFhpa2 virus (\bullet) or control pF2 virus (\square) were incubated (18 h, 37 °C) with sulfate labeled ECM-derived soluble HSPG (peak I). The incubation medium was then subjected to gel filtration on Sepharose 6B. Low molecular weight HS degradation fragments (peak II) were produced only during incubation with the pFhpa2 infected cells, but there was no degradation of the HSPG substrate (\diamond) by lysates of pF2 infected cells.

FIGs. 3a-b demonstrate degradation of soluble sulfate labeled HSPG substrate by the culture medium of pFhpa2 and pFhpa4 infected cells. Culture media of High Five cells infected with pFhpa2 (3a) or pFhpa4 (3b) viruses (\bullet), or with control viruses (\square) were incubated (18 h, 37 °C) with sulfate labeled ECM-derived soluble HSPG (peak I, \diamond). The incubation media were then subjected to gel filtration on Sepharose 6B. Low molecular weight HS degradation fragments (peak II) were produced only during incubation with the hpa gene containing viruses. There was no degradation of the HSPG substrate by the culture medium of cells infected with control viruses.

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FIG. 4 presents size fractionation of heparanase activity expressed by pFhpa2 infected cells. Culture medium of pFhpa2 infected High Five cells was applied onto a 50 kDa cut-off membrane. Heparanase activity (conversion of the peak I substrate, (*) into peak II HS degradation fragments) was found in the high (> 50 kDa) (•), but not low (< 50 kDa) (o) molecular weight compartment.

FIGs. 5a-b demonstrate the effect of heparin on heparanase activity expressed by pFhpa2 and pFhpa4 infected High Five cells. Culture media of pFhpa2 (5a) and pFhpa4 (5b) infected High Five cells were incubated (18 h, 37 °C) with sulfate labeled ECM-derived soluble HSPG (peak I, \Rightarrow) in the absence (\Rightarrow) or presence (\Rightarrow) of 10 µg/ml heparin. Production of low molecular weight HS degradation fragments was completely abolished in the presence of heparin, a potent inhibitor of heparanase activity (6, 7).

FIGs. 6a-b demonstrate degradation of sulfate labeled intact ECM by virus infected High Five and Sf21 cells. High Five (6a) and Sf21 (6b) cells were plated on sulfate labeled ECM and infected (48 h, 28 °C) with pFhpa4 (•) or control pF1 (□) viruses. Control non-infected Sf21 cells (R) were

plated on the labeled ECM as well. The pH of the cultured medium was adjusted to 6.0 - 6.2 followed by 24 h incubation at 37 °C. Sulfate labeled material released into the incubation medium was analyzed by gel filtration on Sepharose 6B. HS degradation fragments were produced only by cells infected with the *hpa* containing virus.

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FIG. 7a-b demonstrate degradation of sulfate labeled intact ECM by virus infected cells. High Five (7a) and Sf21 (7b) cells were plated on sulfate labeled ECM and infected (48 h, 28 °C) with pFhpa4 (•) or control pF1 (□) viruses. Control non-infected Sf21 cells (R) were plate on labeled ECM as well. The pH of the cultured medium was adjusted to 6.0 - 6.2, followed by 48 h incubation at 28 °C. Sulfate labeled degradation fragments released into the incubation medium was analyzed by gel filtration on Sepharose 6B. HS degradation fragments were produced only by cells infected with the hpa containing virus.

FIGs. 8a-b demonstrate degradation of sulfate labeled intact ECM by the culture medium of pFhpa4 infected cells. Culture media of High Five (8a) and Sf21 (8b) cells that were infected with pFhpa4 (•) or control pF1 (□) viruses were incubated (48 h, 37 °C, pH 6.0) with intact sulfate labeled ECM. The ECM was also incubated with the culture medium of control non-infected Sf21 cells (R). Sulfate labeled material released into the reaction mixture was subjected to gel filtration analysis. Heparanase activity was detected only in the culture medium of pFhpa4 infected cells.

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FIGs. 9a-b demonstrate the effect of heparin on heparanase activity in the culture medium of pFhpa4 infected cells. Sulfate labeled ECM was incubated (24 h, 37 °C, pH 6.0) with culture medium of pFhpa4 infected High Five (9a) and Sf21 (9b) cells in the absence (\bullet) or presence (V) of 10 μ g/ml heparin. Sulfate labeled material released into the incubation medium was subjected to gel filtration on Sepharose 6B. Heparanase activity (production of peak II HS degradation fragments) was completely inhibited in the presence of heparin.

FIGs. 10a-b demonstrate purification of recombinant heparanase on heparin-Sepharose. Culture medium of Sf21 cells infected with pFhpa4 virus was subjected to heparin-Sepharose chromatography. Elution of fractions was performed with 0.35 - 2 M NaCl gradient (*). Heparanase activity in the eluted fractions is demonstrated in Figure 10a (•). Fractions 15-28 were subjected to 15 % SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining. A correlation is demonstrated between a

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major protein band (MW \sim 63,000) in fractions 19 - 24 and heparanase activity.

FIGs. 11a-b demonstrate purification of recombinant heparanase on a Superdex 75 gel filtration column. Active fractions eluted from heparin-Sepharose (Figure 10a) were pooled, concentrated and applied onto Superdex 75 FPLC column. Fractions were collected and aliquots of each fraction were tested for heparanase activity (C, Figure 11a) and analyzed by SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining (Figure 11b). A correlation is seen between the appearance of a major protein band (MW \sim 63,000) in fractions 4 - 7 and heparanase activity.

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FIGs. 12a-e demonstrate expression of the hpa gene by RT-PCR with total RNA from human embryonal tissues (12a), human extraembryonal tissues (12b) and cell lines from different origins (12c-e). RT-PCR products using hpa specific primers (I), primers for GAPDH housekeeping gene (II), and control reactions without reverse transcriptase demonstrating absence of genomic DNA or other contamination in RNA samples (III). M- DNA molecular weight marker VI (Boehringer Mannheim). For 12a: lane 1 - neutrophil cells (adult), lane 2 - muscle, lane 3 - thymus, lane 4 - heart, lane 5 - adrenal. For 12b: lane 1 - kidney, lane 2 placenta (8 weeks), lane 3 - placenta (11 weeks), lanes 4-7 - mole (complete hydatidiform mole), lane 8 - cytotrophoblast cells (freshly isolated), lane 9 cytotrophoblast cells (1.5 h in vitro), lane 10 - cytotrophoblast cells (6 h in vitro), lane 11 - cytotrophoblast cells (18 h in vitro), lane 12 cytotrophoblast cells (48 h in vitro). For 12c: lane 1 - JAR bladder cell line, lane 2 - NCITT testicular tumor cell line, lane 3 - SW-480 human hepatoma cell line, lane 4 - HTR (cytotrophoblasts transformed by SV40), lane 5 -HPTLP-I hepatocellular carcinoma cell line, lane 6 - EJ-28 bladder carcinoma cell line. For 12d: lane 1 - SK-hep-1 human hepatoma cell line, lane 2 - DAMI human megakaryocytic cell line, lane 3 - DAMI cell line + PMA, lane 4 - CHRF cell line + PMA, lane 5 - CHRF cell line. For 12e: lane 1 - ABAE bovine aortic endothelial cells, lane 2 - 1063 human ovarian cell line, lane 3 - human breast carcinoma MDA435 cell line, lane 4 human breast carcinoma MDA231 cell line.

FIG. 13 presents a comparison between nucleotide sequences of the human hpa and a mouse EST cDNA fragment (SEQ ID NO:12) which is 80 % homologous to the 3' end (starting at nucleotide 1066 of SEQ ID NO:9) of the human hpa. The aligned termination codons are underlined.

FIG. 14 demonstrates the chromosomal localization of the *hpa* gene. PCR products of DNA derived from somatic cell hybrids and of genomic DNA of hamster, mouse and human of were separated on 0.7 % agarose gel following amplification with *hpa* specific primers. Lane 1 – Lambda DNA digested with *Bst*EII, lane 2 – no DNA control, lanes 3 – 29, PCR amplification products. Lanes 3-5 – human, mouse and hamster genomic DNA, respectively. Lanes 6-29, human monochromosomal somatic cell hybrids representing chromosomes 1-22 and X and Y, respectively. Lane 30 – Lambda DNA digested with *Bst*EII. An amplification product of approximately 2.8 Kb is observed only in lanes 5 and 9, representing human genomic DNA and DNA derived from cell hybrid carrying human chromosome 4, respectively. These results demonstrate that the *hpa* gene is localized in human chromosome 4.

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FIG. 15 demonstrates the genomic exon-intron structure of the human hpa locus (top) and the relative positions of the lambda clones used as sequencing templates to sequence the locus (below). The vertical rectangles represent exons (E) and the horizontal lines therebetween represent introns (I), upstream (U) and downstream (D) regions. Continuous lines represent DNA fragments, which were used for sequence analysis. The discontinuous line in lambda 6 represent a region, which overlaps with lambda 8 and hence was not analyzed. The plasmid contains a PCR product, which bridges the gap between L3 and L6.

FIG. 16 presents the nucleotide sequence of the genomic region of the hpa gene. Exon sequences appear in upper case and intron sequences in lower case. The deduced amino acid sequence of the exons is printed below the nucleotide sequence. Two predicted transcription start sites are shown in bold.

FIG. 17 presents an alignment of the amino acid sequences of human heparanase, mouse and partial sequences of rat homologues. The human and the mouse sequences were determined by sequence analysis of the isolated cDNAs. The rat sequence is derived from two different EST clones, which represent two different regions (5' and 3') of the rat hpa cDNA. The human sequence and the amino acids in the mouse and rat homologues, which are identical to the human sequence, appear in bold.

FIG. 18 presents a heparanase Zoo blot. Ten micrograms of genomic DNA from various sources were digested with *Eco*RI and separated on 0.7 % agarose – TBE gel. Following electrophoresis, the was gel treated with HCl and than with NaOH and the DNA fragments were downward

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transferred to a nylon membrane (Hybond N+, Amersham) with 0.4 N NaOH. The membrane was hybridized with a 1.6 Kb DNA probe that contained the entire *hpa* cDNA. Lane order: H – Human; M – Mouse; Rt – Rat; P – Pig; Cw – Cow; Hr – Horse; S – Sheep; Rb – Rabbit; D – Dog; Ch – Chicken; F – Fish. Size markers (Lambda *BsteII*) are shown on the left

FIG. 19 demonstrates the secondary structure prediction for heparanase performed using the PHD server – Profile network Prediction Heidelberg. H – helix, E – extended (beta strand), The glutamic acid predicted as the proton donor is marked by asterisk and the possible nucleophiles are underlined.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

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The present invention is of a polynucleotide or nucleic acid, referred to hereinbelow interchangeably as hpa, hpa cDNA or hpa gene or identified by its SEQ ID NOs, encoding a polypeptide having heparanase activity, vectors or nucleic acid constructs including same and which are used for over-expression or antisense inhibition of heparanase, genetically modified cells expressing same, recombinant protein having heparanase activity, antisense oligonucleotides and ribozymes for heparanase modulation, and heparanase promoter sequences which can be used to direct the expression of desired genes.

Before explaining at least one embodiment of the invention in detail, it is to be understood that the invention is not limited in its application to the details of construction and the arrangement of the components set forth in the following description or illustrated in the drawings. The invention is capable of other embodiments or of being practiced or carried out in various ways. Also, it is to be understood that the phraseology and terminology employed herein is for the purpose of description and should not be regarded as limiting.

Cloning of the human and mouse hpa genes, cDNAs and genomic sequence (for human), encoding heparanase and expressing recombinant heparanase by transfected cells is reported herein. These are the first mammalian heparanase genes to be cloned.

A purified preparation of heparanase isolated from human hepatoma cells was subjected to tryptic digestion and microsequencing.

The YGPDVGQPR (SEQ ID NO:8) sequence revealed was used to screen EST databases for homology to the corresponding back translated

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DNA sequences. Two closely related EST sequences were identified and were thereafter found to be identical.

Both clones contained an insert of 1020 bp which includes an open reading frame of 973 bp followed by a 3' untranslated region of 27 bp and a Poly A tail, whereas a translation start site was not identified.

Cloning of the missing 5' end was performed by PCR amplification of DNA from placenta Marathon RACE cDNA composite using primers selected according to the EST clones sequence and the linkers of the composite.

A 900 bp PCR fragment, partially overlapping with the identified 3' encoding EST clones was obtained. The joined cDNA fragment (hpa), 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes, as shown in Figure 1 and SEQ ID NO:11, a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons.

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A single nucleotide difference at position 799 (A to T) between the EST clones and the PCR amplified cDNA was observed. This difference results in a single amino acid substitution (Tyr to Phe) (Figure 1). Furthermore, the published EST sequences contained an unidentified nucleotide, which following DNA sequencing of both the EST clones was resolved into two nucleotides (G and C at positions 1630 and 1631 in SEQ ID NO:9, respectively).

The ability of the *hpa* gene product to catalyze degradation of heparan sulfate in an *in vitro* assay was examined by expressing the entire open reading frame in insect cells, using the Baculovirus expression system.

Extracts and conditioned media of cells infected with virus containing the *hpa* gene, demonstrated a high level of heparan sulfate degradation activity both towards soluble ECM-derived HSPG and intact ECM, which was inhibited by heparin, while cells infected with a similar construct containing no *hpa* gene had no such activity, nor did non-infected cells.

The expression pattern of hpa RNA in various tissues and cell lines was investigated using RT-PCR. It was found to be expressed only in tissues and cells previously known to have heparanase activity.

Cloning an extended 5' sequence was enabled from the human SK-hep1 cell line by PCR amplification using the Marathon RACE. The 5' extended sequence of the SK-hep1 hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQ ID NO:9).

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The assembled sequence contained an open reading frame, SEQ ID NOs: 13 and 15, which encodes, as shown in SEQ ID NOs:14 and 15, a polypeptide of 592 amino acids, with a calculated molecular weight of 66,407 daltons. This open reading frame was shown to direct the expression of catalytically active heparanase in a mammalian cell expression system. The expressed heparanase was detectable by anti heparanase antibodies in Western blot analysis.

A panel of monochromosomal human/CHO and human/mouse somatic cell hybrids was used to localize the human heparanase gene to human chromosome 4. The newly isolated heparanase sequence can therefore be used to identify a chromosome region harboring a human heparanase gene in a chromosome spread.

. The hpa cDNA was then used as a probe to screen a a human genomic library. Several phages were positive. These phages were analyzed and were found to cover most of the hpa locus, except for a small portion which was recovered by bridging PCR. The hpa locus covers about 50,000 bp. The hpa gene includes 12 exons separated by 11 introns.

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RT-PCR performed on a variety of cells revealed alternatively spliced hpa transcripts.

The amino acid sequence of human heparanase was used to search for homologous sequences in the DNA and protein databases. Several human EST's were identified, as well as mouse sequences highly homologous to human heparanase. The following mouse EST's were identified AA177901, AA674378, AA67997, AA047943, AA690179, AI122034, all sharing an identical sequence and correspond to amino acids 336-543 of the human heparanase sequence. The entire mouse heparanase cDNA was cloned, based on the nucleotide sequence of the mouse EST's using Marathon cDNA libraries. The mouse and the human hpa genes share an average homology of 78 % between the nucleotide sequences and 81 % similarity between the deduced amino acid sequences. hpa homologous sequences from rat were also uncovered (EST's AI060284 and AI237828).

Homology search of heparanase amino acid sequence against the DNA and the protein databases and prediction of its protein secondary structure enabled to identify candidate amino acids that participate in the heparanase active site.

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Expression of hpa antisense in mammalian cell lines resulted in about five fold decrease in the number of recoverable cells as compared to controls.

Human *Hpa* cDNA was shown to hybridize with genomic DNAs of a variety of mammalian species and with an avian.

The human and mouse *hpa* promoters were identified and the human promoter was tested positive in directing the expression of a reporter gene.

Thus, according to the present invention there is provided an isolated nucleic acid comprising a genomic, complementary or composite polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

The phrase "composite polynucleotide sequence" refers to a sequence which includes exonal sequences required to encode the polypeptide having heparanase activity, as well as any number of intronal sequences. The intronal sequences can be of any source and typically will include conserved splicing signal sequences. Such intronal sequences may further include cis acting expression regulatory elements.

The term "heparanase catalytic activity" or its equivalent term "heparanase activity" both refer to a mammalian endoglycosidase hydrolyzing activity which is specific for heparan or heparan sulfate proteoglycan substrates, as opposed to the activity of bacterial enzymes (heparinase I, II and III) which degrade heparin or heparan sulfate by means of β -elimination (37).

According to a preferred embodiment of the present invention the polynucleotide or a portion thereof is hybridizable with SEQ ID NOs: 9, 13, 42, 43 or a portion thereof at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 µg/ml salmon sperm DNA, and ³²p labeled probe and wash at 68 °C with 3, 2, 1, 0.5 or 0.1 x SSC and 0.1 % SDS.

According to another preferred embodiment of the present invention the polynucleotide or a portion thereof is at least 60 %, preferably at least 65 %, more preferably at least 70 %, more preferably at least 75 %, more preferably at least 80 %, more preferably at least 85 %, more preferably at least 90 %, most preferably, 95-100 % identical with SEQ ID NOs: 9, 13, 42, 43 or portions thereof as determined using the Bestfit procedure of the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the university of Wisconsin (gap creation penalty - 12, gap extension penalty - 4 - which are the default parameters).

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According to another preferred embodiment of the present invention the polypeptide encoded by the polynucleotide sequence is as set forth in SEQ ID NOs:10, 14, 44 or portions thereof having heparanase catalytic activity. Such portions are expected to include amino acids Asp-Glu 224-225 (SEQ ID NO:10), which can serve as proton donors and glutamic acid 343 or 396 which can serve as a nucleophile.

According to another preferred embodiment of the present invention the polypeptide encoded by the polynucleotide sequence is at least 60 %, preferably at least 65 %, more preferably at least 70 %, more preferably at least 85 %, more preferably at least 80 %, more preferably at least 85 %, more preferably at least 90 %, most preferably, 95-100 % homologous (both similar and identical acids) to SEQ ID NOs:10, 14, 44 or portions thereof as determined with the Smith-Waterman algorithm, using the Bioaccelerator platform developed by Compugene (gapop: 10.0, gapext: 0.5, matrix: blosum62, see also the description to Figure 17).

Further according to the present invention there is provided a nucleic acid construct comprising the isolated nucleic acid described herein. The construct may and preferably further include an origin of replication and

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The construct or vector can be of any type. It may be a phage which infects bacteria or a virus which infects eukaryotic cells. It may also be a plasmid, phagemid, cosmid, bacmid or an artificial chromosome.

Further according to the present invention there is provided a host cell comprising the nucleic acid construct described herein. The host cell can be of any type. It may be a prokaryotic cell, an eukaryotic cell, a cell line, or a cell as a portion of an organism. The polynucleotide encoding heparanase can be permanently or transiently present in the cell. In other words, genetically modified cells obtained following stable or transient transfection, transformation or transduction are all within the scope of the present invention. The polynucleotide can be present in the cell in low copy (say 1-5 copies) or high copy number (say 5-50 copies or more). It may be integrated in one or more chromosomes at any location or be present as an extrachromosomal material.

The present invention is further directed at providing a heparanase over-expression system which includes a cell overexpressing heparanase catalytic activity. The cell may be a genetically modified host cell transiently or stably transfected or transformed with any suitable vector which includes a polynucleotide sequence encoding a polypeptide having

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heparanase activity and a suitable promoter and enhancer sequences to direct over-expression of heparanase. However, the overexpressing cell may also be a product of an insertion (e.g., via homologous recombination) of a promoter and/or enhancer sequence downstream to the endogenous heparanase gene of the expressing cell, which will direct over-expression from the endogenous gene.

The term "over-expression" as used herein in the specification and claims below refers to a level of expression which is higher than a basal level of expression typically characterizing a given cell under otherwise identical conditions.

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According to another aspect the present invention provides an antisense oligonucleotide comprising a polynucleotide or a polynucleotide analog of at least 10, preferably 11-15, more preferably 16-17, more preferably 18, more preferably 19-25, more preferably 26-35, most preferably 35-100 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity. The antisense oligonucleotide can be used for downregulating heparanase activity by *in vivo* administration thereof to a patient. As such, the antisense oligonucleotide according to the present invention can be used to treat types of cancers which are characterized by impaired (over) expression of heparanase, and are dependent on the expression of heparanase for proliferating or forming metastases.

The antisense oligonucleotide can be DNA or RNA or even include nucleotide analogs, examples of which are provided in the Background section hereinabove. The antisense oligonucleotide according to the present invention can be synthetic and is preferably prepared by solid phase synthesis. In addition, it can be of any desired length which still provides specific base pairing (e.g., 8 or 10, preferably more, nucleotides long) and it can include mismatches that do not hamper base pairing under physiological conditions.

Further according to the present invention there is provided a pharmaceutical composition comprising the antisense oligonucleotide herein described and a pharmaceutically acceptable carrier. The carrier can be, for example, a liposome loadable with the antisense oligonucleotide.

According to a preferred embodiment of the present invention the antisense oligonucleotide further includes a ribozyme sequence. The ribozyme sequence serves to cleave a heparanase RNA molecule to which

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the antisense oligonucleotide binds, to thereby downregulate heparanase expression.

Further according to the present invention there is provided an antisense nucleic acid construct comprising a promoter sequence and a polynucleotide sequence directing the synthesis of an antisense RNA sequence of at least 10 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity. Like the antisense oligonucleotide, the antisense construct can be used for downregulating heparanase activity by *in vivo* administration thereof to a patient. As such, the antisense construct, like the antisense oligonucleotide, according to the present invention can be used to treat types of cancers which are characterized by impaired (over) expression of heparanase, and are dependent on the expression of heparanase for proliferating or forming metastases.

Thus, further according to the present invention there is provided a pharmaceutical composition comprising the antisense construct herein described and a pharmaceutically acceptable carrier. The carrier can be, for example, a liposome loadable with the antisense construct.

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Formulations for topical administration may include, but are not limited to, lotions, ointments, gels, creams, suppositories, drops, liquids, sprays and powders. Conventional pharmaceutical carriers, aqueous, powder or oily bases, thickeners and the like may be necessary or desirable. Coated condoms, stents, active pads, and other medical devices may also be useful. Compositions for oral administration include powders or granules, suspensions or solutions in water or non-aqueous media, sachets, capsules or tablets. Thickeners, diluents, flavorings, dispersing aids, emulsifiers or binders may be desirable. Formulations for parenteral administration may include, but are not limited to, sterile aqueous solutions which may also contain buffers, diluents and other suitable additives.

Dosing is dependent on severity and responsiveness of the condition to be treated, but will normally be one or more doses per day, week or month with course of treatment lasting from several days to several months or until a cure is effected or a diminution of disease state is achieved. Persons ordinarily skilled in the art can easily determine optimum dosages, dosing methodologies and repetition rates.

Further according to the present invention there is provided a nucleic acid construct comprising a polynucleotide sequence functioning as a promoter, the polynucleotide sequence is derived from SEQ ID NO:42 and

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includes at least nucleotides 2135-2635, preferably 2235-2635, more preferably 2335-2635, more preferably 2435-2635, most preferably 2535-2635 thereof, or SEQ ID NO:43 and includes at least nucleotides 1-420, preferably 120-420, more preferably 220-420, most preferably 320-420, thereof. These nucleotides are shown in the example section that follows to direct the synthesis of a reporter gene in transformed cells. Thus, further according to the present invention there is provided a method of expressing a polynucleotide sequence comprising the step of ligating the polynucleotide sequence downstream to either of the promoter sequences described herein. Heparanase promoters can be isolated from a variety of mammalian an other species by cloning genomic regions present 5' to the coding sequence thereof. This can be readily achievable by one ordinarily skilled in the art using the heparanase polynucleotides described herein, which are shown in the Examples section that follows to participate in efficient cross species hybridization.

Further according to the present invention there is provided a recombinant protein comprising a polypeptide having heparanase catalytic activity. The protein according to the present invention include modifications known as post translational modifications, including, but not limited to, proteolysis (e.g., removal of a signal peptide and of a pro- or preprotein sequence), methionine modification, glycosylation, alkylation (e.g., methylation), acetylation, etc. According to preferred embodiments the polypeptide includes at least a portion of SEQ ID NOs:10, 14 or 44, the portion has heparanase catalytic activity. According to preferred embodiments of the present invention the protein is encoded by any of the above described isolated nucleic acids. Further according to the present invention there is provided a pharmaceutical composition comprising, as an active ingredient, the recombinant protein described herein.

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The recombinant protein may be purified by any conventional protein purification procedure close to homogeneity and/or be mixed with additives. The recombinant protein may be manufactured using any of the genetically modified cells described above, which include any of the expression nucleic acid constructs described herein. The recombinant protein may be in any form. It may be in a crystallized form, a dehydrated powder form or in solution. The recombinant protein may be useful in obtaining pure heparanase, which in turn may be useful in eliciting antiheparanase antibodies, either poly or monoclonal antibodies, and as a

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screening active ingredient in an anti-heparanase inhibitors or drugs screening assay or system.

Further according to the present invention there is provided a method of identifying a chromosome region harboring a human heparanase gene in a chromosome spread. the method is executed implementing the following method steps, in which in a first step the chromosome spread (either interphase or metaphase spread) is hybridized with a tagged polynucleotide probe encoding heparanase. The tag is preferably a fluorescent tag. In a second step according to the method the chromosome spread is washed, thereby excess of non-hybridized probe is removed. Finally, signals associated with the hybridized tagged polynucleotide probe are searched for, wherein detected signals being indicative of a chromosome region harboring the human heparanase gene. One ordinarily skilled in the art would know how to use the sequences disclosed herein in suitable labeling reactions and how to use the tagged probes to detect, using *in situ* hybridization, a chromosome region harboring a human heparanase gene.

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Further according to the present invention there is provided a method of *in vivo* eliciting anti-heparanase antibodies comprising the steps of administering a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*. Accordingly, there is provided also a DNA vaccine for *in vivo* eliciting anti-heparanase antibodies comprising a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*. The vaccine optionally further includes a pharmaceutically acceptable carrier, such as a virus, liposome or an antigen presenting cell. Alternatively, the vaccine is employed as a naked DNA vaccine

The present invention can be used to develop treatments for various diseases, to develop diagnostic assays for these diseases and to provide new tools for basic research especially in the fields of medicine and biology.

Specifically, the present invention can be used to develop new drugs to inhibit tumor cell metastasis, inflammation and autoimmunity. The identification of the *hpa* gene encoding for the heparanase enzyme enables the production of a recombinant enzyme in heterologous expression systems.

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Furthermore, the present invention can be used to modulate bioavailability of heparin-binding growth factors, cellular responses to heparin-binding growth factors (e.g., bFGF, VEGF) and cytokines (e.g., IL-8), cell interaction with plasma lipoproteins, cellular susceptibility to viral, protozoa and some bacterial infections, and disintegration of neurodegenerative plaques. Recombinant heparanase offers a potential treatment for wound healing, angiogenesis, restenosis, atherosclerosis, inflammation, neurodegenerative diseases (such as, for example, Genstmann-Straussler Syndrome, Creutzfeldt-Jakob disease, Scrape and Alzheimer's disease) and certain viral and some bacterial and protozoa infections. Recombinant heparanase can be used to neutralize plasma heparin, as a potential replacement of protamine.

As used herein, the term "modulate" includes substantially inhibiting, slowing or reversing the progression of a disease, substantially ameliorating clinical symptoms of a disease or condition, or substantially preventing the appearance of clinical symptoms of a disease or condition. A "modulator" therefore includes an agent which may modulate a disease or condition. Modulation of viral, protozoa and bacterial infections includes any effect which substantially interrupts, prevents or reduces any viral, bacterial or protozoa activity and/or stage of the virus, bacterium or protozoon life cycle, or which reduces or prevents infection by the virus, bacterium or protozoon in a subject, such as a human or lower animal.

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As used herein, the term "wound" includes any injury to any portion of the body of a subject including, but not limited to, acute conditions such as thermal burns, chemical burns, radiation burns, burns caused by excess exposure to ultraviolet radiation such as sunburn, damage to bodily tissues such as the perineum as a result of labor and childbirth, including injuries sustained during medical procedures such as episiotomies, trauma-induced injuries including cuts, those injuries sustained in automobile and other mechanical accidents, and those caused by bullets, knives and other weapons, and post-surgical injuries, as well as chronic conditions such as pressure sores, bedsores, conditions related to diabetes and poor circulation, and all types of acne, etc.

Anti-heparanase antibodies, raised against the recombinant enzyme, would be useful for immunodetection and diagnosis of micrometastases, autoimmune lesions and renal failure in biopsy specimens, plasma samples, and body fluids. Such antibodies may also serve as neutralizing agents for heparanase activity.

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The genomic heparanase sequences described herein can be used to construct knock-in and knock-out constructs. Such constructs include a fragment of 10-20 Kb of a heparanase locus and a negative and a positive selection markers and can be used to provide heparanase knock-in and knock-out animal models by methods known to the skilled artisan. Such animal models can be used for studying the function of heparanase in developmental processes, and in normal as well as pathological processes. They can also serve as an experimental model for testing drugs and gene therapy protocols. The complementary heparanase sequence (cDNA) can be used to derive transgenic animals, overexpressing heparanase for same. Alternatively, if cloned in the antisense orientation, the complementary heparanase sequence (cDNA) can be used to derive transgenic animals under-expressing heparanase for same.

The heparanase promoter sequences described herein and other cis regulatory elements linked to the heparanase locus can be used to regulated the expression of genes. For example, these promoters can be used to direct the expression of a cytotoxic protein, such as TNF, in tumor cells. It will be appreciated that heparanase itself is abnormally expressed under the control of its own promoter and other cis acting elements in a variety of tumors, and its expression is correlated with metastasis. It is also abnormally highly expressed in inflammatory cells. The introns of the heparanase gene can be used for the same purpose, as it is known that introns, especially upstream introns include cis acting element which affect expression. A heparanase promoter fused to a reporter protein can be used to study/monitor its activity.

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The polynucleotide sequences described herein can also be used to provide DNA vaccines which will elicit in vivo anti heparanase antibodies. Such vaccines can therefore be used to combat inflammatory and cancer.

Antisense oligonucleotides derived according to the heparanase sequences described herein, especially such oligonucleotides supplemented with ribozyme activity, can be used to modulate heparanase expression. Such oligonucleotides can be from the coding region, from the introns or promoter specific. Antisense heparanase nucleic acid constructs can similarly function, as well known in the art.

The heparanase sequences described herein can be used to study the catalytic mechanism of heparanase. Carefully selected site directed mutagenesis can be employed to provide modified heparanase proteins

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having modified characteristics in terms of, for example, substrate specificity, sensitivity to inhibitors, etc.

While studying heparanase expression in a variety of cell types alternatively spliced transcripts were identified. Such transcripts if found characteristic of certain pathological conditions can be used as markers for such conditions. Such transcripts are expected to direct the synthesis of heparanases with altered functions.

Additional objects, advantages, and novel features of the present invention will become apparent to one ordinarily skilled in the art upon examination of the following examples, which are not intended to be limiting. Additionally, each of the various embodiments and aspects of the present invention as delineated hereinabove and as claimed in the claims section below finds experimental support in the following examples.

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EXAMPLES

Generally, the nomenclature used herein and the laboratory procedures in recombinant DNA technology described below are those well known and commonly employed in the art. Standard techniques are used for cloning, DNA and RNA isolation, amplification and purification. Generally enzymatic reactions involving DNA ligase, DNA polymerase, restriction endonucleases and the like are performed according to the manufacturers' specifications. These techniques and various other techniques are generally performed according to Sambrook et al., Molecular Cloning--A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. (1989), which is incorporated herein by reference. Other general references are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the convenience of the reader. All the information contained therein is incorporated herein by reference.

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The following protocols and experimental details are referenced in the Examples that follow:

Purification and characterization of heparanase from a human hepatoma cell line and human placenta: A human hepatoma cell line (Skhep-1) was chosen as a source for purification of a human tumor-derived heparanase. Purification was essentially as described in U.S. Pat. No. 5,362,641 to Fuks, which is incorporated by reference as if fully set forth

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herein. Briefly, 500 liter, $5x10^{11}$ cells were grown in suspension and the heparanase enzyme was purified about 240,000 fold by applying the following steps: (i) cation exchange (CM-Sephadex) chromatography performed at pH 6.0, 0.3-1.4 M NaCl gradient; (ii) cation exchange (CM-Sephadex) chromatography performed at pH 7.4 in the presence of 0.1% CHAPS, 0.3-1.1 M NaCl gradient; (iii) heparin-Sepharose chromatography performed at pH 7.4 in the presence of 0.1% CHAPS, 0.35-1.1 M NaCl gradient; (iv) ConA-Sepharose chromatography performed at pH 6.0 in buffer containing 0.1 % CHAPS and 1 M NaCl, elution with 0.25 M α -methyl mannoside; and (v) HPLC cation exchange (Mono-S) chromatography performed at pH 7.4 in the presence of 0.1 % CHAPS, 0.25-1 M NaCl gradient.

Active fractions were pooled, precipitated with TCA and the precipitate subjected to SDS polyacrylamide gel electrophoresis and/or tryptic digestion and reverse phase HPLC. Tryptic peptides of the purified protein were separated by reverse phase HPLC (C8 column) and homogeneous peaks were subjected to amino acid sequence analysis.

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The purified enzyme was applied to reverse phase HPLC and subjected to N-terminal amino acid sequencing using the amino acid sequencer (Applied Biosystems).

Cells: Cultures of bovine corneal endothelial cells (BCECs) were established from steer eyes as previously described (19, 38). Stock cultures were maintained in DMEM (1 g glucose/liter) supplemented with 10 % newborn calf serum and 5 % FCS. bFGF (1 ng/ml) was added every other day during the phase of active cell growth (13, 14).

Preparation of dishes coated with ECM: BCECs (second to fifth passage) were plated into 4-well plates at an initial density of 2 x 10⁵ cells/ml, and cultured in sulfate-free Fisher medium plus 5 % dextran T-40 for 12 days. Na₂³⁵SO₄ (25 μCi/ml) was added on day 1 and 5 after seeding and the cultures were incubated with the label without medium change. The subendothelial ECM was exposed by dissolving (5 min., room temperature) the cell layer with PBS containing 0.5 % Triton X-100 and 20 mM NH₄OH, followed by four washes with PBS. The ECM remained intact, free of cellular debris and firmly attached to the entire area of the tissue culture dish (19, 22).

To prepare soluble sulfate labeled proteoglycans (peak I material), the ECM was digested with trypsin (25 μ g/ml, 6 h, 37 °C), the digest was concentrated by reverse dialysis and the concentrated material was applied

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onto a Sepharose 6B gel filtration column. The resulting high molecular weight material (Kav< 0.2, peak I) was collected. More than 80 % of the labeled material was shown to be composed of heparan sulfate proteoglycans (11, 39).

Heparanase activity: Cells (1 x 106/35-mm dish), cell lysates or conditioned media were incubated on top of 35S-labeled ECM (18 h, 37 °C) in the presence of 20 mM phosphate buffer (pH 6.2). Cell lysates and conditioned media were also incubated with sulfate labeled peak I material (10-20 µl). The incubation medium was collected, centrifuged (18,000 x g, 4 °C, 3 min.), and sulfate labeled material analyzed by gel filtration on a Sepharose CL-6B column (0.9 x 30 cm). Fractions (0.2 ml) were eluted with PBS at a flow rate of 5 ml/h and counted for radioactivity using Biofluor scintillation fluid. The excluded volume (Vo) was marked by blue dextran and the total included volume (Vt) by phenol red. The latter was shown to comigrate with free sulfate (7, 11, 23). Degradation fragments of HS side chains were eluted from Sepharose 6B at 0.5 < Kav < 0.8 (peak II) (7, 11, 23). A nearly intact HSPG released from ECM by trypsin - and, to a lower extent, during incubation with PBS alone - was eluted next to Vo (Kav < 0.2, peak I). Recoveries of labeled material applied on the columns ranged from 85 to 95 % in different experiments (11). Each experiment was performed at least three times and the variation of elution positions (Kav values) did not exceed +/- 15 %.

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Cloning of hpa cDNA: cDNA clones 257548 and 260138 were obtained from the I.M.A.G.E Consortium (2130 Memorial Parkway SW, Hunstville, AL 35801). The cDNAs were originally cloned in EcoRI and NotI cloning sites in the plasmid vector pT3T7D-Pac. Although these clones are reported to be somewhat different, DNA sequencing demonstrated that these clones are identical to one another. Marathon RACE (rapid amplification of cDNA ends) human placenta (poly-A) cDNA composite was a gift of Prof. Yossi Shiloh of Tel Aviv University. This composite is vector free, as it includes reverse transcribed cDNA fragments to which double, partially single stranded adapters are attached on both sides. The construction of the specific composite employed is described in reference 39a.

Amplification of hp3 PCR fragment was performed according to the protocol provided by Clontech laboratories. The template used for amplification was a sample taken from the above composite. The primers used for amplification were:

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First step: 5'-primer: AP1: 5'-CCATCCTAATACGACTCACT ATAGGGC-3', SEQ ID NO:1; 3'-primer: HPL229: 5'-GTAGTGATGCCA TGTAACTGAATC-3', SEQ ID NO:2.

Second step: nested 5'-primer: AP2: 5'-ACTCACTATAGGGCTCG AGCGGC-3', SEQ ID NO:3; nested 3'- primer: HPL171: 5'-GCATCTTAGCCGTCTTTCTTCG-3', SEQ ID NO:4. The HPL229 and HPL171 were selected according to the sequence of the EST clones. They include nucleotides 933-956 and 876-897 of SEQ ID NO:9, respectively.

PCR program was 94 °C - 4 min., followed by 30 cycles of 94 °C - 40 sec., 62 °C - 1 min., 72 °C - 2.5 min. Amplification was performed with Expand High Fidelity (Boehringer Mannheim). The resulting ca. 900 bp hp3 PCR product was digested with *BfrI* and *PvuII*. Clone 257548 (phpa1) was digested with *EcoRI*, followed by end filling and was then further digested with *BfrI*. Thereafter the *PvuII* - *BfrI* fragment of the hp3 PCR product was cloned into the blunt end - *BfrI* end of clone phpa1 which resulted in having the entire cDNA cloned in pT3T7-pac vector, designated phpa2.

RT-PCR: RNA was prepared using TRI-Reagent (Molecular research center Inc.) according to the manufacturer instructions. 1.25 µg were taken for reverse transcription reaction using MuMLV Reverse transcriptase (Gibco BRL) and Oligo (dT)₁₅ primer, SEQ ID NO:5, (Promega). Amplification of the resultant first strand cDNA was performed with *Taq* polymerase (Promega). The following primers were used:

25 HPU-355: 5'-TTCGATCCCAAGAAGGAATCAAC-3', SEQ ID NO:6, nucleotides 372-394 in SEQ ID NOs:9 or 11.
HPL-229: 5'-GTAGTGATGCCATGTAACTGAATC-3', SEQ ID NO:7, nucleotides 933-956 in SEQ ID NOs:9 or 11.

PCR program: 94 °C - 4 min., followed by 30 cycles of 94 °C - 40 sec., 62 °C - 1 min., 72 °C - 1 min.

Alternatively, total RNA was prepared from cell cultures using Trireagent (Molecular Research Center, Inc.) according to the manufacturer recommendation. Poly A+ RNA was isolated from total RNA using mRNA separator (Clontech). Reverse transcription was performed with total RNA using Superscript II (GibcoBRL). PCR was performed with Expand high fidelity (Boehringer Mannheim). Primers used for amplification were as follows:

Hpu-685, 5'-GAGCAGCCAGGTGAGCCCAAGAT-3', SEQ ID NO:24

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Hpu-355, 5'-TTCGATCCCAAGAAGGAATCAAC-3', SEQ ID NO:25

Hpu 565, 5'-AGCTCTGTAGATGTGCTATACAC-3', SEQ ID NO:26

Hpl 967, 5'-TCAGATGCAAGCAGCAACTTTGGC-3', SEQ ID NO:27 Hpl 171, 5'-GCATCTTAGCCGTCTTTCTTCG-3', SEO ID NO:28

Hpl 229, 5'-GTAGTGATGCCATGTAACTGAATC-3', SEQ ID NO:29

PCR reaction was performed as follows: 94 °C 3 minutes, followed by 32 cycles of 94 °C 40 seconds, 64 °C 1 minute, 72 °C 3 minutes, and one cycle 72 °C, 7 minutes.

Expression of recombinant heparanase in insect cells: Cells, High Five and Sf21 insect cell lines were maintained as monolayer cultures in SF900II-SFM medium (GibcoBRL).

Recombinant Baculovirus: Recombinant virus containing the hpa gene was constructed using the Bac to Bac system (GibcoBRL). The transfer vector pFastBac was digested with SalI and NotI and ligated with a 1.7 kb fragment of phpa2 digested with XhoI and NotI. The resulting plasmid was designated pFasthpa2. An identical plasmid designated pFasthpa4 was prepared as a duplicate and both independently served for further experimentations. Recombinant bacmid was generated according to the instructions of the manufacturer with pFasthpa2, pFasthpa4 and with pFastBac. The latter served as a negative control. Recombinant bacmid DNAs were transfected into Sf21 insect cells. Five days after transfection recombinant viruses were harvested and used to infect High Five insect cells, 3 x 106 cells in T-25 flasks. Cells were harvested 2 - 3 days after infection. 4 x 106 cells were centrifuged and resuspended in a reaction buffer containing 20 mM phosphate citrate buffer, 50 mM NaCl. Cells underwent three cycles of freeze and thaw and lysates were stored at -80 °C. Conditioned medium was stored at 4 °C.

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Partial purification of recombinant heparanase: Partial purification of recombinant heparanase was performed by heparin-Sepharose column chromatography followed by Superdex 75 column gel filtration. Culture medium (150 ml) of Sf21 cells infected with pFhpa4 virus was subjected to heparin-Sepharose chromatography. Elution of 1 ml fractions was performed with 0.35 - 2 M NaCl gradient in presence of 0.1 % CHAPS and 1 mM DTT in 10 mM sodium acetate buffer, pH 5.0. A 25 μ l sample of each fraction was tested for heparanase activity. Heparanase activity was eluted at the range of 0.65 - 1.1 M NaCl (fractions 18-26, Figure 10a). 5 μ l of each fraction was subjected to 15 % SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining.

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Active fractions eluted from heparin-Sepharose (Figure 10a) were pooled and concentrated (x 6) on YM3 cut-off membrane. 0.5 ml of the concentrated material was applied onto 30 ml Superdex 75 FPLC column equilibrated with 10 mM sodium acetate buffer, pH 5.0, containing 0.8 M NaCl, 1 mM DTT and 0.1 % CHAPS. Fractions (0.56 ml) were collected at a flow rate of 0.75 ml/min. Aliquots of each fraction were tested for heparanase activity and were subjected to SDS-polyacrylamide gel electrophoresis followed by silver nitrate staining (Figure 11b).

PCR amplification of genomic DNA: 94 °C 3 minutes, followed by 32 cycles of 94 °C 45 seconds, 64 °C 1 minute, 68 °C 5 minutes, and one cycle at 72 °C, 7 minutes. Primers used for amplification of genomic DNA included:

GHpu-L3 5'-AGGCACCCTAGAGATGTTCCAG-3', SEQ ID NO:30 GHpl-L6 5'-GAAGATTTCTGTTTCCATGACGTG-3', SEQ ID NO:31.

Screening of genomic libraries: A human genomic library in Lambda phage EMBLE3 SP6/T7 (Clontech, Paulo Alto, CA) was screened. 5 x 10⁵ plaques were plated at 5 x 10⁴ pfu/plate on NZCYM agar/top agarose plates. Phages were absorbed on nylon membranes in duplicates (Qiagen). Hybridization was performed at 65 °C in 5 x SSC, 5 x Denhart's, 10 % dextran sulfate, 100 μg/ml Salmon sperm, ³²p labeled probe (10⁶ cpm/ml). A 1.6 kb fragment, containing the entire hpa cDNA was labeled by random priming (Boehringer Mannheim). Following hybridization membranes were washed once with 2 x SSC, 0.1 % SDS at 65 °C for 20 minutes, and twice with 0.2 x SSC, 0.1 % SDS at 65 °C for 15 minutes. Hybridizing plaques were picked, and plated at 100 pfu/plate. Hybridization was performed as above and single isolated positive plaques were picked.

Phage DNA was extracted using a Lambda DNA extraction kit (Qiagen). DNA was digested with *XhoI* and *EcoRI*, separated on 0.7 % agarose gel and transferred to nylon membrane Hybond N+ (Amersham). Hybridization and washes were performed as above.

cDNA Sequence analysis: Sequence determinations were performed with vector specific and gene specific primers, using an automated DNA sequencer (Applied Biosystems, model 373A). Each nucleotide was read from at least two independent primers.

Genomic sequence analysis: Large-scale sequencing was performed by Commonwealth Biotechnology Incorporation.

Isolation of mouse hpa: Mouse hpa cDNA was amplified from either Marathon ready cDNA library of mouse embryo or from mRNA isolated from mouse melanoma cell line BL6, using the Marathon RACE kit from Clontech. Both procedures were performed according to the manufacturer's recommendation.

Primers used for PCR amplification of mouse hpa:

Mhpl773 5'-CCACACTGAATGTAATACTGAAGTG-3', SEQ ID NO:32

MHpl736 5'-CGAAGCTCTGGAACTCGGCAAG-3', SEQ ID NO:33

MHpl83 5'-GCCAGCTGCAAAGGTGTTGGAC-3', SEQ ID NO:34

Mhpl152 5'-AACACCTGCCTCATCACGACTTC-3', SEQ ID NO:35

Mhpl114 5'-GCCAGGCTGGCGTCGATGGTGA-3', SEQ ID NO:36

MHpl103 5'-GTCGATGGTGATGGACAGGAAC-3', SEQ ID NO:37

Ap1 5'-GTAATACGACTCACTATAGGGC-3', SEQ ID NO:38 (Genome walker)

15 Ap2 5'-ACTATAGGGCACGCGTGGT-3', SEQ ID NO:39 (Genome walker)
Ap1 5'-CCATCCTAATACGACTCACTATAGGGC-3', SEQ ID NO:40 (Marathon RACE)
Ap2 5'-ACTCACTATAGGGCTCGAGCGGC-3', SEQ ID NO:41 20 (Marathon RACE)

Southern analysis of genomic DNA: Genomic DNA was extracted from animal or from human blood using Blood and cell culture DNA maxi kit (Qiagene). DNA was digested with *EcoRI*, separated by gel electrophoresis and transferred to a nylon membrane Hybond N+(Amersham). Hybridization was performed at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 μg/ml salmon sperm DNA, and ³²p labeled probe. A 1.6 kb fragment, containing the entire *hpa* cDNA was used as a probe. Following hybridization, the membrane was washed with 3 x SSC, 0.1 % SDS, at 68 °C and exposed to X-ray film for 3 days. Membranes were then washed with 1 x SSC, 0.1 % SDS, at 68 °C and were reexposed for 5 days.

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Construction of hpa promoter-GFP expression vector: Lambda DNA of phage L3, was digested with SacI and BgIII, resulting in a 1712 bp fragment which contained the hpa promoter (877-2688 of SEQ ID NO:42). The pEGFP-1 plasmid (Clontech) was digested with BgIII and SacI and ligated with the 1712 bp fragment of the hpa promoter sequence. The resulting plasmid was designated phpEGL. A second hpa promoter-GFP plasmid was constructed containing a shorter fragment of the hpa promoter

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region: phpEGL was digested with *Hind*III, and the resulting 1095 bp fragment (nucleotides 1593-2688 of SEQ ID NO:42) was ligated with *Hind*III digested pEGFP-1. The resulting plasmid was designated phpEGS.

Computer analysis of sequences: Homology searches were performed using several computer servers, and various databases. Blast 2.0 service, at the NCBI server was used to screen the protein database swplus and DNA databases such as GenBank, EMBL, and the EST databases. Blast 2.0 search was performed using the basic search option of the NCBI server. Sequence analysis and alignments were done using the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the university of Wisconsin. Alignments of two sequences were performed using Bestfit (gap creation penalty - 12, gap extension penalty - 4). Protein homology search was performed with the Smith-Waterman algorithm, using the Bioaccelerator platform developed by Compugene. The protein database swplus was searched using the following parameters: gapop: 10.0, gapext: 0.5, matrix: blosum62. Blocks homology was performed using the Blocks WWW server developed at Fred Hutchinson Cancer Research Center in Seattle, Washington, USA. Secondary structure prediction was performed using the PHD server -Profile network Prediction Heidelberg. Fold recognition (threading) was performed using the UCLA-DOE structure prediction server. The method used for prediction was gonnet+predss. Alignment of three sequences was performed using the pileup application (gap creation penalty - 5, gap extension penalty - 1). Promoter analysis was performed using TSSW and TSSG programs (BCM Search Launcher Human Genome Center, Baylor College of Medicine, Houston TX).

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EXAMPLE 1

Cloning of human hpa cDNA

Purified fraction of heparanase isolated from human hepatoma cells (SK-hep-1) was subjected to tryptic digestion and microsequencing. EST (Expressed Sequence Tag) databases were screened for homology to the back translated DNA sequences corresponding to the obtained peptides. Two EST sequences (accession Nos. N41349 and N45367) contained a DNA sequence encoding the peptide YGPDVGQPR (SEQ ID NO:8). These two sequences were derived from clones 257548 and 260138 (I.M.A.G.E Consortium) prepared from 8 to 9 weeks placenta cDNA library (Soares). Both clones which were found to be identical contained an insert

of 1020 bp which included an open reading frame (ORF) of 973 bp followed by a 3' untranslated region of 27 bp and a Poly A tail. No translation start site (AUG) was identified at the 5' end of these clones.

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Cloning of the missing 5' end was performed by PCR amplification of DNA from a placenta Marathon RACE cDNA composite. A 900 bp fragment (designated hp3), partially overlapping with the identified 3' encoding EST clones was obtained.

The joined cDNA fragment, 1721 bp long (SEQ ID NO:9), contained an open reading frame which encodes, as shown in Figure 1 and SEQ ID NO:11, a polypeptide of 543 amino acids (SEQ ID NO:10) with a calculated molecular weight of 61,192 daltons. The 3' end of the partial cDNA inserts contained in clones 257548 and 260138 started at nucleotide G⁷²¹ of SEQ ID NO:9 and Figure 1.

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As further shown in Figure 1, there was a single sequence discrepancy between the EST clones and the PCR amplified sequence, which led to an amino acid substitution from Tyr²⁴⁶ in the EST to Phe²⁴⁶ in the amplified cDNA. The nucleotide sequence of the PCR amplified cDNA fragment was verified from two independent amplification products. The new gene was designated *hpa*.

As stated above, the 3' end of the partial cDNA inserts contained in EST clones 257548 and 260138 started at nucleotide 721 of hpa (SEQ ID NO:9). The ability of the hpa cDNA to form stable secondary structures, such as stem and loop structures involving nucleotide stretches in the vicinity of position 721 was investigated using computer modeling. It was found that stable stem and loop structures are likely to be formed involving nucleotides 698-724 (SEQ ID NO:9). In addition, a high GC content, up to 70 %, characterizes the 5' end region of the hpa gene, as compared to about only 40 % in the 3' region. These findings may explain the immature termination and therefore lack of 5' ends in the EST clones.

To examine the ability of the hpa gene product to catalyze degradation of heparan sulfate in an in vitro assay the entire open reading frame was expressed in insect cells, using the Baculovirus expression system. Extracts of cells, infected with virus containing the hpa gene, demonstrated a high level of heparan sulfate degradation activity, while cells infected with a similar construct containing no hpa gene had no such activity, nor did non-infected cells. These results are further demonstrated in the following Examples.

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EXAMPLE 2

Degradation of soluble ECM-derived HSPG

Monolayer cultures of High Five cells were infected (72 h, 28 °C) with recombinant Bacoluvirus containing the pFasthpa plasmid or with control virus containing an insert free plasmid. The cells were harvested and lysed in heparanase reaction buffer by three cycles of freezing and thawing. The cell lysates were then incubated (18 h, 37 °C) with sulfate labeled, ECM-derived HSPG (peak I), followed by gel filtration analysis (Sepharose 6B) of the reaction mixture.

As shown in Figure 2, the substrate alone included almost entirely high molecular weight (Mr) material eluted next to V_0 (peak I, fractions 5-20, Kav < 0.35). A similar elution pattern was obtained when the HSPG substrate was incubated with lysates of cells that were infected with control virus. In contrast, incubation of the HSPG substrate with lysates of cells infected with the *hpa* containing virus resulted in a complete conversion of the high Mr substrate into low Mr labeled degradation fragments (peak II, fractions 22-35, 0.5 < Kav < 0.75).

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Fragments eluted in peak II were shown to be degradation products of heparan sulfate, as they were (i) 5- to 6-fold smaller than intact heparan sulfate side chains (Kav approx. 0.33) released from ECM by treatment with either alkaline borohydride or papain; and (ii) resistant to further digestion with papain or chondroitinase ABC, and susceptible to deamination by nitrous acid (6, 11). Similar results (not shown) were obtained with Sf21 cells. Again, heparanase activity was detected in cells infected with the *hpa* containing virus (pFhpa), but not with control virus (pF). This result was obtained with two independently generated recombinant viruses. Lysates of control not infected High Five cells failed to degrade the HSPG substrate.

In subsequent experiments, the labeled HSPG substrate was incubated with medium conditioned by infected High Five or Sf21 cells.

As shown in Figures 3a-b, heparanase activity, reflected by the conversion of the high Mr peak I substrate into the low Mr peak II which represents HS degradation fragments, was found in the culture medium of cells infected with the pFhpa2 or pFhpa4 viruses, but not with the control pF1 or pF2 viruses. No heparanase activity was detected in the culture medium of control non-infected High Five or Sf21 cells.

The medium of cells infected with the pFhpa4 virus was passed through a 50 kDa cut off membrane to obtain a crude estimation of the

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molecular weight of the recombinant heparanase enzyme. As demonstrated in Figure 4, all the enzymatic activity was retained in the upper compartment and there was no activity in the flow through (<50 kDa) material. This result is consistent with the expected molecular weight of the hpa gene product.

In order to further characterize the *hpa* product the inhibitory effect of heparin, a potent inhibitor of heparanase mediated HS degradation (40) was examined.

As demonstrated in Figures 5a-b, conversion of the peak I substrate into peak II HS degradation fragments was completely abolished in the presence of heparin.

Altogether, these results indicate that the heparanase enzyme is expressed in an active form by insect cells infected with Baculovirus containing the newly identified human hpa gene.

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EXAMPLE 3

Degradation of HSPG in intact ECM

Next, the ability of intact infected insect cells to degrade HS in intact, naturally produced ECM was investigated. For this purpose, High Five or Sf21 cells were seeded on metabolically sulfate labeled ECM followed by infection (48 h, 28 °C) with either the pFhpa4 or control pF2 viruses. The pH of the medium was then adjusted to pH 6.2-6.4 and the cells further incubated with the labeled ECM for another 48 h at 28 °C or 24 h at 37 °C. Sulfate labeled material released into the incubation medium was analyzed by gel filtration on Sepharose 6B.

As shown in Figures 6a-b and 7a-b, incubation of the ECM with cells infected with the control pF2 virus resulted in a constant release of labeled material that consisted almost entirely (>90%) of high Mr fragments (peak I) eluted with or next to V_0 . It was previously shown that a proteolytic activity residing in the ECM itself and/or expressed by cells is responsible for release of the high Mr material (6). This nearly intact HSPG provides a soluble substrate for subsequent degradation by heparanase, as also indicated by the relatively large amount of peak I material accumulating when the heparanase enzyme is inhibited by heparin (6, 7, 12, Figure 9). On the other hand, incubation of the labeled ECM with cells infected with the pFhpa4 virus resulted in release of 60-70% of the ECM-associated radioactivity in the form of low Mr sulfate-labeled fragments (peak II, 0.5 <Kav< 0.75), regardless of whether the infected cells were incubated with

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the ECM at 28 °C or 37 °C. Control intact non-infected Sf21 or High Five cells failed to degrade the ECM HS side chains.

In subsequent experiments, as demonstrated in Figures 8a-b, High Five and Sf21 cells were infected (96 h, 28 °C) with pFhpa4 or control pF1 viruses and the culture medium incubated with sulfate-labeled ECM. Low Mr HS degradation fragments were released from the ECM only upon incubation with medium conditioned by pFhpa4 infected cells. As shown in Figure 9, production of these fragments was abolished in the presence of heparin. No heparanase activity was detected in the culture medium of control, non-infected cells. These results indicate that the heparanase enzyme expressed by cells infected with the pFhpa4 virus is capable of degrading HS when complexed to other macromolecular constituents (i.e. fibronectin, laminin, collagen) of a naturally produced intact ECM, in a manner similar to that reported for highly metastatic tumor cells or activated cells of the immune system (6, 7).

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EXAMPLE 4

Purification of recombinant human heparanase

The recombinant heparanase was partially purified from medium of pFhpa4 infected Sf21 cells by Heparin-Sepharose chromatography (Figure 10a) followed by gel filtration of the pooled active fractions over an FPLC Superdex 75 column (Figure 11a). A \sim 63 kDa protein was observed, whose quantity, as was detected by silver stained SDS-polyacrylamide gel electrophoresis, correlated with heparanase activity in the relevant column fractions (Figures 10b and 11b, respectively). This protein was not detected in the culture medium of cells infected with the control pF1 virus and was subjected to a similar fractionation on heparin-Sepharose (not shown).

EXAMPLE 5

30 Expression of the human hpa cDNA in various cell types, organs and tissues

Referring now to Figures 12a-e, RT-PCR was applied to evaluate the expression of the *hpa* gene by various cell types and tissues. For this purpose, total RNA was reverse transcribed and amplified. The expected 585 bp long cDNA was clearly demonstrated in human kidney, placenta (8 and 11 weeks) and mole tissues, as well as in freshly isolated and short termed (1.5-48 h) cultured human placental cytotrophoblastic cells (Figure 12a), all known to express a high heparanase activity (41). The *hpa*

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transcript was also expressed by normal human neutrophils (Figure 12b). In contrast, there was no detectable expression of the *hpa* mRNA in embryonic human muscle tissue, thymus, heart and adrenal (Figure 12b). The *hpa* gene was expressed by several, but not all, human bladder carcinoma cell lines (Figure 12c), SK hepatoma (SK-hep-1), ovarian carcinoma (OV 1063), breast carcinoma (435, 231), melanoma and megakaryocytic (DAMI, CHRF) human cell lines (Figures 12d-e).

The above described expression pattern of the *hpa* transcript was determined to be in a very good correlation with heparanase activity levels determined in various tissues and cell types (not shown).

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EXAMPLE 6

Isolation of an extended 5' end of hpa cDNA from human SK-hep1 cell line

The 5' end of hpa cDNA was isolated from human SK-hep1 cell line by PCR amplification using the Marathon RACE (rapid amplification of cDNA ends) kit (Clontech). Total RNA was prepared from SK-hep1 cells using the TRI-Reagent (Molecular research center Inc.) according to the manufacturer instructions. Poly A+ RNA was isolated using the mRNA separator kit (Clonetech).

The Marahton RACE SK-hep1 cDNA composite was constructed according to the manufacturer recommendations. First round of amplification was performed using an adaptor specific primer AP1: 5'-CCATCCTAATACG ACTCACTATAGGGC-3', SEQ ID NO:1, and a hpa specific antisense primer hpl-629: 5'-CCCCAGGAGCAGCAGCATCAG-3', SEQ ID NO:17, corresponding to nucleotides 119-99 of SEQ ID NO:9. The resulting PCR product was subjected to a second round of amplification using adaptor specific nested primer AP2: ACTCACTATAGGGCTCGAGCGGC-3', SEQ ID NO:3, and a hpa specific hpl-666 5'antisense nested primer AGGCTTCGAGCGCAGCAGCAT-3', SEQ ID NO:18, corresponding to nucleotides 83-63 of SEQ ID NO:9. The PCR program was as follows: a hot start of 94 °C for 1 minute, followed by 30 cycles of 90 °C - 30 seconds, 68 °C - 4 minutes. The resulting 300 bp DNA fragment was extracted from an agarose gel and cloned into the vector pGEM-T Easy (Promega). The resulting recombinant plasmid was designated pHPSK1.

The nucleotide sequence of the pHPSK1 insert was determined and it was found to contain 62 nucleotides of the 5' end of the placenta hpa cDNA

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(SEQ ID NO:9) and additional 178 nucleotides upstream, the first 178 nucleotides of SEQ ID NOs:13 and 15.

A single nucleotide discrepancy was identified between the SK-hepl cDNA and the placenta cDNA. The "T" derivative at position 9 of the placenta cDNA (SEQ ID NO:9), is replaced by a "C" derivative at the corresponding position 187 of the SK-hepl cDNA (SEQ ID NO:13).

The discrepancy is likely to be due to a mutation at the 5' end of the placenta cDNA clone as confirmed by sequence analysis of sevsral additional cDNA clones isolated from placenta, which like the SK-hep1 cDNA contained C at position 9 of SEQ ID NO:9.

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The 5' extended sequence of the SK-hep1 hpa cDNA was assembled with the sequence of the hpa cDNA isolated from human placenta (SEQ ID NO:9). The assembled sequence contained an open reading frame which encodes, as shown in SEQ ID NOs:14 and 15, a polypeptide of 592 amino acids with a calculated molecular weight of 66,407 daltons. The open reading frame is flanked by 93 bp 5' untranslated region (UTR).

EXAMPLE 7

Isolation of the upstream genomic region of the hpa gene

The upstream region of the *hpa* gene was isolated using the Genome Walker kit (Clontech) according to the manufacturer recommendations. The kit includes five human genomic DNA samples each digested with a different restriction endonuclease creating blunt ends: *Eco*RV, *Sca*I, *Dra*I, *Pvu*II and *Ssp*I.

The blunt ended DNA fragments are ligated to partially single stranded adaptors. The Genomic DNA samples were subjected to PCR amplification using the adaptor specific primer and a gene specific primer. Amplification was performed with Expand High Fidelity (Boehringer Mannheim).

A first round of amplification was performed using the ap1 primer: 5'-G TAATACGACTCACTATAGGGC-3', SEQ ID NO:19, and the hpa specific antisense primer hpl-666: 5'-AGGCTTCGAGCGCAGCAGCAT-3', SEQ ID NO:18, corresponding to nucleotides 83 – 63 of SEQ ID NO:9. The PCR program was as follows: a hot start of 94 °C - 3 minutes, followed by 36 cycles of 94 °C - 40 seconds, 67 °C - 4 minutes.

The PCR products of the first amplification were diluted 1:50. One μ l of the diluted sample was used as a template for a second amplification using a nested adaptor specific primer ap2: 5'-

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ACTATAGGGCACGCGTGGT-3', SEQ ID NO:20, and a hpa specific antisense primer hpl-690, 5'-CTTGGGCTCACC TGGCTGCTC-3', SEQ ID NO:21, corresponding to nucleotides 62-42 of SEQ ID NO:9. The resulting amplification products were analyzed using agarose gel electrophoresis. Five different PCR products were obtained from the five amplification reactions. A DNA fragment of approximately 750 bp which was obtained from the SspI digested DNA sample was gel extracted. The purified fragment was ligated into the plasmid vector pGEM-T Easy (Promega). The resulting recombinant plasmid was designated pGHP6905 and the nucleotide sequence of the hpa insert was determined.

A partial sequence of 594 nucleotides is shown in SEQ ID NO:16. The last nucleotide in SEQ ID NO:13 corresponds to nucleotide 93 in SEQ ID:13. The DNA sequence in SEQ ID NO:16 contains the 5' region of the hpa cDNA and 501 nucleotides of the genomic upstream region which are predicted to contain the promoter region of the hpa gene.

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EXAMPLE 8

Expression of the 592 amino acids HPA polypeptide in a human 293 cell line

The 592 amino acids open reading frame (SEQ ID NOs:13 and 15) was constructed by ligation of the 110 bp corresponding to the 5' end of the SK-hep1 hpa cDNA with the placenta cDNA. More specifically the Marathon RACE - PCR amplification product of the placenta hpa DNA was digested with SacI and an approximately 1 kb fragment was ligated into a SacI-digested pGHP6905 plasmid. The resulting plasmid was digested with Earl and AatII. The EarI sticky ends were blunted and an approximately 280 bp EarI/blunt-AatII fragment was isolated. This fragment was ligated with pFasthpa digested with EcoRI which was blunt ended using Klenow fragment and further digested with AatII. The resulting plasmid contained a 1827 bp insert which includes an open reading frame of 1776 bp, 31 bp of 3' UTR and 21 bp of 5' UTR. This plasmid was designated pFastLhpa.

A mammalian expression vector was constructed to drive the expression of the 592 amino acids heparanase polypeptide in human cells. The hpa cDNA was excised prom pFastLhpa with BssHII and NotI. The resulting 1850 bp BssHII-NotI fragment was ligated to a mammalian expression vector pSI (Promega) digested with MluI and NotI. The resulting recombinant plasmid, pSIhpaMet2 was transfected into a human 293 embryonic kidney cell line.

Transient expression of the 592 amino-acids heparanase was examined by western blot analysis and the enzymatic activity was tested using the gel shift assay. Both these procedures are described in length in U.S. Pat. application No. 09/071,739, filed May 1, 1998, which is incorporated by reference as if fully set forth herein. Cells were harvested 3 days following transfection. Harvested cells were re-suspended in lysis buffer containing 150 mM NaCl, 50 mM Tris pH 7.5, 1% Triton X-100, 1 mM PMSF and protease inhibitor cocktail (Boehringer Mannheim). 40 µg protein extract samples were used for separation on a SDS-PAGE. Proteins were transferred onto a PVDF Hybond-P membrane (Amersham). The membrane was incubated with an affinity purified polyclonal anti heparanase antibody, as described in U.S. Pat. application No. 09/071,739. A major band of approximately 50 kDa was observed in the transfected cells as well as a minor band of approximately 65 kDa. A similar pattern was observed in extracts of cells transfected with the pShpa as demonstrated in U.S. Pat. application No. 09/071,739. These two bands probably represent two forms of the recombinant heparanase protein produced by the transfected cells. The 65 kDa protein probably represents a heparanase precursor, while the 50 kDa protein is suggested herein to be the processed or mature form.

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The catalytic activity of the recombinant protein expressed in the pShpaMet2 transfected cells was tested by gel shift assay. Cell extracts of transfected and of mock transfected cells were incubated overnight with heparin (6 µg in each reaction) at 37 °C, in the presence of 20 mM phosphate citrate buffer pH 5.4, 1 mM CaCl₂, 1 mM DTT and 50 mM NaCl. Reaction mixtures were then separated on a 10 % polyacrylamide gel. The catalytic activity of the recombinant heparanase was clearly demonstrated by a faster migration of the heparin molecules incubated with the transfected cell extract as compared to the control. Faster migration indicates the disappearance of high molecular weight heparin molecules and the generation of low molecular weight degradation products.

EXAMPLE 9

Chromosomal localization of the hpa gene

Chromosomal mapping of the *hpa* gene was performed utilizing a panel of monochromosomal human/CHO and human/mouse somatic cell hybrids, obtained from the UK HGMP Resource Center (Cambridge, England).

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40 ng of each of the somatic cell hybrid DNA samples were subjected to PCR amplification using the *hpa* primers: hpu565 5'-AGCTCTGTAGATGTGC TATACAC-3', SEQ ID NO:22, corresponding to nucleotides 564-586 of SEQ ID NO:9 and an antisense primer hpl171 5'-GCATCTTAGCCGTCTTTCTTCG-3', SEQ ID NO:23, corresponding to nucleotides 897-876 of SEQ ID NO:9.

The PCR program was as follows: a hot start of 94 °C - 3 minutes, followed by 7 cycles of 94 °C - 45 seconds, 66 °C - 1 minute, 68 °C - 5 minutes, followed by 30 cycles of 94 °C - 45 seconds, 62 °C - 1 minute, 68 °C - 5 minutes, and a 10 minutes final extension at 72 °C.

The reactions were performed with Expand long PCR (Boehringer Mannheim). The resulting amplification products were analyzed using agarose gel electrophoresis. As demonstrated in Figure 14, a single band of approximately 2.8 Kb was obtained from chromosome 4, as well as from the control human genomic DNA. A 2.8 kb amplification product is expected based on amplification of the genomic *hpa* clone (data not shown). No amplification products were obtained neither in the control DNA samples of hamster and mouse nor in somatic hybrids of other human chromosome.

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EXAMPLE 10

Human genomic clone encoding heparanase

Five plaques were isolated following screening of a human genomic library and were designated L3-1, L5-1, L8-1, L10-1 and L6-1. The phage DNAs were analyzed by Southern hybridization and by PCR with hpa specific and vector specific primers. Southern analysis was performed with three fragments of hpa cDNA: a PvuII-BamHI fragment (nucleotides 32-450, SEQ ID NO:9), a BamHI-NdeI fragment (nucleotides 451-1102, SEQ ID NO:9) and an NdeI-XhoI fragment (nucleotides 1103-1721, SEQ ID NO:9).

Following Southern analysis, phages L3, L6, L8 were selected for further analysis. A scheme of the genomic region and the relative position of the three phage clones is depicted in Figure 15. A 2 kb DNA fragment containing the gap between phages L6 and L3 was PCR amplified from human genomic DNA with two gene specific primers GHpuL3 and GHplL6. The PCR product was cloned into the plasmid vector pGEM-Teasy (Promega).

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Large scale DNA sequencing of the three Lambda clones and the amplified fragment was performed with Lambda purified DNA by primer walking. A nucleotide sequence of 44,898 bp was analyzed (Figure 16, SEQ ID NO:42). Comparison of the genomic sequence with that of hpa cDNA revealed 12 exons separated by 11 introns (Figures 15 an 16). The genomic organization of the hpa gene is depicted in Figure 15 (top). The sequence include the coding region from the first ATG to the stop codon which spans 39,113 nucleotides, 2742 nucleotides upstream of the first ATG and 3043 nucleotides downstream of the stop codon. Splice site consensus sequences were identified at exon/intron junctions.

EXAMPLE 11

Alternative splicing

Several minor RT-PCR products were obtained from various cell types, following amplification with hpa specific primers. Each one found to contain a deletion of one or two exons. Some of these PCR products contain ORFs, which encode potential shorter proteins.

Table 1 below summarizes the alternative spliced products isolated from various cell lines.

Fragments of similar sizes were obtained following amplification with two cell lines, placenta and platelets.

	Cell type	Nucleotides deleted	Exons deleted	ORF
25	Platelets	1047-1267	8, 9	+
	Platelets	1154-1267	9	-
	Platelets	289-435, 562-735	2, 4	-
	Sk-hep1, platelets, Zr75	562-735	4	+
	Sk-hep1 (hepatoma)	561-904	4, 5	-
30	Zr75 (breast carcinoma)	96-203	1 (partial)	+

EXAMPLE 12

Mouse and rat hpa

EST databases were screened for sequences homologous to the *hpa* gene. Three mouse EST's were identified (accession No. Aa177901, from mouse spleen, Aa067997 from mouse skin, Aa47943 from mouse embryo), assembled into a 824 bp cDNA fragment which contains a partial open reading frame (lacking a 5' end) of 629 bp and a 3' untranslated region of

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195 bp (SEQ ID NO:12). As shown in Figure 13, the coding region is 80 % similar to the 3' end of the hpa cDNA sequence. These EST's are probably cDNA fragments of the mouse hpa homolog that encodes for the mouse heparanase.

Searching for consensus protein domains revealed an amino terminal homology between the heparanase and several precursor proteins such as Procollagen Alpha 1 precursor, Tyrosine-protein kinase-RYK, Fibulin-1, Insulin-like growth factor binding protein and several others. The amino terminus is highly hydrophobic and contains a potential trans-membrane domain. The homology to known signal peptide sequences suggests that it could function as a signal peptide for protein localization.

The amino acid sequence of human heparanase was used to search for homologous sequences in the DNA and protein databases. Several human EST's were identified, as well as mouse sequences highly homologous to human heparanase. The following mouse EST's were identified AA177901, AA674378, AA67997, AA047943, AA690179, AI122034, all sharing an identical sequence and correspond to amino acids 336-543 of the human heparanase sequence. The entire mouse heparanase cDNA was cloned, based on the nucleotide sequence of the mouse EST's. PCR primers were designed and a Marathon RACE was performed using a Marathon cDNA library from 15 days mouse embryo (Clontech) and from BL6 mouse melanoma cell line. The mouse hpa homologous cDNA was isolated following several amplification steps. A 1.1 kb fragment was amplified from mouse embryo Marathon cDNA library. The first cycle of amplification was performed with primers mhpl773 and Ap1 and the second cycle with primers mhp1736 and AP2. A 1.1 kb fragment was then amplified from BL6 Marathon cDNA library. The first cycle of amplification was performed with the primers mhpl152 and Ap1, and the second with mhpl83 and AP2. The combined sequence was homologous to nucleotides 157 - 1702 of the human hpa cDNA, which encode amino acids 33-543. The 5' end of the mouse hpa gene was isolated from a mouse genomic DNA library using the Genome Walker kit (Clontech). An 0.9 kb fragment was amplified from a DraI digested Genome walker DNA library. The first cycle of amplification was performed with primers mhpl114 and Apl and the second with primers mhpl103 and AP2. The assembled sequence (SEQ ID NOs:43, 45) is 2396 nucleotides long. It contains an open reading frame of 1605 nucleotides, which encode a polypeptide of 535 amino acids (SEQ ID NOs:44, 45), 196 nucleotides of 3' untranslated

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region (UTR), and anupstream sequence which includes the promoter region and the 5'-UTR of the mouse hpa cDNA.. According to two promoter predicting programs TSSW and TSSG, the transcription start site is localized to nucleotide 431 of SEQ ID NOs:43, 45, 163 nucleotides upstream of the first ATG codon. The 431 upstream genomic sequence contains the promoter region. A TATA box is predicted at position 394 of SEQ ID NOs:43, 45. The mouse and the human hpa genes share an average homology of 78 % between the nucleotide sequences and 81 % similarity between the deduced amino acid sequences.

Search for hpa homologous sequences, using the Blast 2.0 server revealed two EST's from rat: AI060284 (385 nucleotides, SEQ ID NO:46) which is homologous to the amino terminus (68 % similarity to amino acids 12-136) of human heparanase and AI237828 (541 nucleotides, SEQ ID NO:47) which is homologous to the carboxyl terminus (81 % similarity to amino acids 500-543) of human heparanase, and contains a 3'-UTR. A comparison between the human heparanase and the mouse and rat homologous sequences is demonstrated in Figure 17.

EXAMPLE 13

Prediction of heparanase active site

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Homology search of heparanase amino acid sequence against the DNA and the protein databases revealed no significant homologies. The protein secondary structure as predicted by the PHD program consists of alternating alpha helices and beta sheets. The fold recognition server of UCLA predicted alpha/beta barrel structure, with under-threshold confidence.

Five of 15 proteins, which were predicted to have most similar folds, were glycosyl hydrolases from various organisms: 1xyza – xylanase from Clostridium Thermocellum, 1pbga – 6-phospho-beta-δ-galactosidase from Lactococcus Lactis, 1amy – alpha-amylase from Barley, 1ecea – endocellulase from Acidothermus Cellulolyticus and 1qbc – hexosaminidase alpha chain, glycosyl hydrolase.

Protein homology search using the bioaccelerator pulled out several proteins, including glycosyl hydrolyses such as beta-fructofuranosidase from *Vicia faba* (broad bean) and from potato, lactase phlorizin hydrolase from human, xylanases from *Clostridium thermocellum* and from *Streptomyces halstedii* and cellulase from *Clostridium thermocellum*. Blocks 9.3 database pulled out the active site of glycosyl hydrolases family

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five, which includes cellulases from various bacteria and fungi. Similar active site motif is shared by several lysosomal acid hydrolases (63) and other glycosyl hydrolases. The common mechanism shared by these enzymes involves two glutamic acid residues, a proton donor and a nucleophile.

Despite the lack of an overall homology between the heparanase and other glycosyl hydolases, the amino acid couple Asp-Glu (NE), which is characteristic of the proton donor of glycosyl hydrolyses of the GH-A clan, was found at positions 224-225 of the human heparanase protein sequence. As in other clan members, this NE couple is located at the end of a β sheet.

Considering the relative location of the proton donor and the predicted secondary structure, the glutamic acid that functions as nucleophile is most likely located at position 343, or at position 396. Identification of the active site and the amino acids directly involved in hydrolysis opens the way for expression of the defined catalytic domain. In addition, it will provide the tools for rational design of enzyme activity either by modification of the microenviroment or catalytic site itself.

EXAMPLE 14

Expression of hpa antisense in mammalian cell lines

A mammalian expression vector Hpa2Kepcdna3 was constructed in order to express *hpa* antisense in mammalian cells. *hpa* cDNA (1.7 kb *Eco*RI fragment) was cloned into the plasmid pCDNA3 in 3'>5' (antisense) orientation. The construct was used to transfect MBT2-T50 and T24P cell lines. 2 x 10⁵ cells in 35 mm plates were transfected using the Fugene protocol (Boehringer Mannheim). 48 hours after transfection cells were trypsinized and seeded in six well plates. 24 hours later G418 was added to initiate selection. The number of colonies per 35 mm plate following 3 weeks:

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	Antisense	No insert	
T24P	15	60	
MBT-T50	1	6	

The lower number of colonies obtained after transfection with hpa antisense, as compared with the control plasmid suggests that the introduction of hpa antisense interfere with cell growth. This experiment demonstrates the use of complementary antisense hpa DNA sequence to

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control heparanase expression in cells. This approach may be used to inhibit expression of heparanase *in vivo*, in, for example, cancer cells and in other pathological processes in which heparanase is involved.

EXAMPLE 15

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Zoo blot

Hpa cDNA was used as a probe to detect homologous sequences in human DNA and in DNA of various animals. The autoradiogram of the Southern analysis is presented in Figure 18. Several bands were detected in human DNA, which correlated with the accepted pattern according to the genomic hpa sequence. Several intense bands were detected in all mammals, while faint bands were detected in chicken. This correlates with the phylogenetic relation between human and the tested animals. intense bands indicate that hpa is conserved among mammals as well as in more genetically distant organisms. The multiple bands patterns suggest that in all animals, like in human, the hpa locus occupy large genomic Alternatively, the various bands could represent homologous sequences and suggest the existence of a gene family, which can be isolated based on their homology to the human hpa reported herein. conservation was actually found, between the isolated human hpa cDNA and the mouse homologue.

EXAMPLE 16

Characterization of the hpa promoter

The DNA sequence upstream of the *hpa* first ATG was subjected to computational analysis in order to localize the predicted transcription start site and to identify potential transcription factors binding sites. Recognition of human PolII promoter region and start of transcription were predicted using the TSSW and TSSG programs. Both programs identified a promoter region upstream of the coding region. TSSW pointed at nucleotide 2644 and TSSG at 2635 of SEQ ID NO:42. These two predicted transcription start sites are located 4 and 13 nucleotides upstream of the longest *hpa* cDNA isolated by RACE.

A hpa promoter-GFP reporter vector was constructed in order to investigate the regulation of hpa transcription. Two constructs were made, containing 1.8 kb and 1.1 kb of the hpa promoter region. The reporter vector was transfected into T50-mouse bladder carcinoma cells. Cells transfected with both constructs exhibited green fluorescence, which

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indicated the promoter activity of the genomic sequence upstream of the *hpa*-coding region. This reporter vector, enables the monitoring of hpa promoter activity, at various conditions and in different cell types and to characterize the factors involved regulation of *hpa* expression.

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Although the invention has been described in conjunction with specific embodiments thereof, it is evident that many alternatives, modifications and variations will be apparent to those skilled in the art. Accordingly, it is intended to embrace all such alternatives, modifications and variations that fall within the spirit and broad scope of the appended claims.

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PCT/US00/03542

WHAT IS CLAIMED IS:

WO 00/52178

1. An isolated nucleic acid comprising a genomic, complementary or composite polynucleotide sequence encoding a polypeptide having heparanase catalytic activity.

- 2. The isolated nucleic acid of claim 1, wherein said polynucleotide or a portion thereof is hybridizable with SEQ ID NOs: 9, 13, 42, 43 or a portion thereof at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 μg/ml salmon sperm DNA, and ³²p labeled probe and wash at 68 °C with 3 x SSC and 0.1 % SDS.
- 3. The isolated nucleic acid of claim 1, wherein said polynucleotide or a portion thereof is at least 60 % identical with SEQ ID NOs: 9, 13, 42, 43 or portions thereof as determined using the Bestfit procedure of the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the university of Wisconsin (gap creation penalty 12, gap extension penalty 4).
- 4. The isolated nucleic acid of claim 1, wherein said polypeptide is as set forth in SEQ ID NOs:10, 14, 44 or portions thereof.
- 5. The isolated nucleic acid of claim 1, wherein said polypeptide is at least 60 % homologous to SEQ ID NOs:10, 14, 44 or portions thereof as determined with the Smith-Waterman algorithm, using the Bioaccelerator platform developed by Compugene (gapop: 10.0, gapext: 0.5, matrix: blosum62).
- 6. A nucleic acid construct comprising the isolated nucleic acid of claim 1.
 - 7. A host cell comprising the nucleic acid construct of claim 6.
- 8. An antisense oligonucleotide comprising a polynucleotide or a polynucleotide analog of at least 10 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity.

- 9. The antisense oligonucleotide of claim 8, wherein said polynucleotide strand encoding said polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 9, 13, 42, or 43.
- 10. The antisense oligonucleotide of claim 8, wherein said polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 10, 14 and 44.
- 11. A method of *in vivo* downregulating heparanase activity comprising the step of *in vivo* administering the antisense oligonucleotide of claim 8.
- 12. A pharmaceutical composition comprising the antisense oligonucleotide of claim 8 and a pharmaceutically acceptable carrier.
- 13. A ribozyme comprising the antisense oligonucleotide of claim 8 and a ribozyme sequence.
- 14. An antisense nucleic acid construct comprising a promoter sequence and a polynucleotide sequence directing the synthesis of an antisense RNA sequence of at least 10 bases being hybridizable *in vivo*, under physiological conditions, with a portion of a polynucleotide strand encoding a polypeptide having heparanase catalytic activity.
- 15. The antisense nucleic acid construct of claim 14, wherein said polynucleotide strand encoding said polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 9, 13, 42 or 43.
- 16. The antisense nucleic acid construct of claim 14, wherein said polypeptide having heparanase catalytic activity is as set forth in SEQ ID NOs: 10, 14 or 44.
- 17. A method of *in vivo* downregulating heparanase activity comprising the step of *in vivo* administering the antisense nucleic acid construct of claim 14.

18. A pharmaceutical composition comprising the antisense nucleic acid construct of claim 14 and a pharmaceutically acceptable carrier.

- 19. A nucleic acid construct comprising a polynucleotide sequence functioning as a promoter, said polynucleotide sequence is derived from SEQ ID NO:42 and includes at least nucleotides 2535-2635 thereof or from SEQ ID NO:43 and includes at least nucleotides 320-420.
- 20. A method of expressing a polynucleotide sequence comprising the step of ligating the polynucleotide sequence into the nucleic acid construct of claim 19, downstream of said polynucleotide sequence derived from SEQ ID NOs:42 or 43.
- 21. A recombinant protein comprising a polypeptide having heparanase catalytic activity.
- 22. The recombinant protein of claim 21, wherein said polypeptide includes at least a portion of SEQ ID NOs:10, 14 or 44.
- 23. The recombinant protein of claim 21, wherein the protein is encoded by a polynucleotide hybridizable with SEQ ID NOs: 9, 13, 42, 43 or a portion thereof at 68 °C in 6 x SSC, 1 % SDS, 5 x Denharts, 10 % dextran sulfate, 100 μ g/ml salmon sperm DNA, and ³²p labeled probe and wash at 68 °C with 3 x SSC and 0.1 % SDS.
- 24. The recombinant protein of claim 21, wherein the protein is encoded by a polynucleotide at least 60 % identical with SEQ ID NOs: 9, 13, 42, 43 or portions thereof as determined using the Bestfit procedure of the DNA sequence analysis software package developed by the Genetic Computer Group (GCG) at the university of Wisconsin (gap creation penalty 12, gap extension penalty 4).
- 25. A pharmaceutical composition comprising, as an active ingredient, the recombinant protein of claim 21.
- 26. A method of identifying a chromosome region harboring a heparanase gene in a chromosome spread comprising the steps of:

- (a) hybridizing the chromosome spread with a tagged polynucleotide probe encoding heparanase;
- (b) washing the chromosome spread, thereby removing excess of non-hybridized probe; and
- (c) searching for signals associated with said hybridized tagged polynucleotide probe, wherein detected signals being indicative of a chromosome region harboring a heparanase gene.
- 27. A method of *in vivo* eliciting anti-heparanase antibodies comprising the steps of administering a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*.
- 28. A DNA vaccine for *in vivo* eliciting anti-heparanase antibodies comprising a nucleic acid construct including a polynucleotide segment corresponding to at least a portion of SEQ ID NOs:9, 13 or 43 and a promoter for directing the expression of said polynucleotide segment *in vivo*.

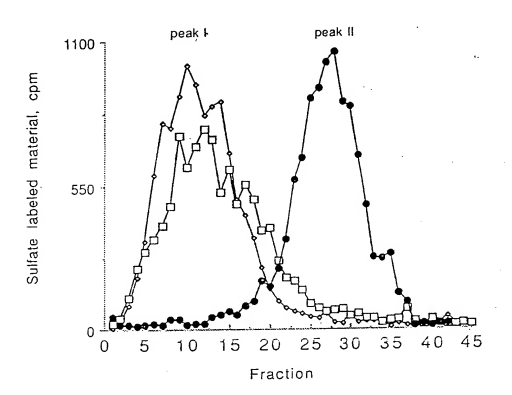
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I D A N L A T D P R P L I L L G S P K L 301 TTOSTACCTTGGCCAGAGGCTTGTCTCCTGCGTACCTGAGGTTTGGTGGCACCAAGACAG RTLARGLSPAYLRFGGTKTĎ 361 ACTTCCTAATTTTCGATCCCAAGAAGGAATCAACCTTTGAAGAGAGAAGTTACTGGCAAT FLIFDPKKESTFBERSTWQS 421 CTCAAGTCAACCAGGATATTTGCAAATATGGATCCATCCCTCCTGATGTGGAGGAGAGT Q V N Q D I C K Y G. S I P P D V B B K L 481 TACGGTTGGAATGGCCCTACCAGGAGCAATTGCTACTCCGAGAACACTACCAGAAAAAGT LBWPYQEQLLLRBHYQKK 541 TCAAGAACAGCACCTACTCAAGAAGCTCTGTAGATGTGCTATACACTTTTGCAAACTGCT KNSTYSRSSVDVLYTFANCS 601 CAGGACTGGACTTGATCTTTGGCCTAAATGCGTTATTAAGAACAGCAGATTTGCAGTGGA G L D L I F G L N A L L R T A D L Q W N 661 ACAGTTCTAATGCTCAGTTGCTCCTGGACTACTGCTCTTCCAAGGGGTATAACATTTCTT S S N A Q L L L D Y C S S K G Y N I S W 721 GGGAACTAGGCAATGAACCTAACAGTTTCCTTAAGAAGGCTGATATTTTCATCAATGGGT ELGNEPNSFLKKADIFINGS TS: CSCASTTAGGAGAAGATTATATTCAATTGCATAAACTTCTAAGAAAGTCCACCTTCAAAA
Q L G E D Y 1 Q L H K L L R K S T F K N
(F) ATGCAAAACTCTATGGTCCTGATGTTGGTCAGCCTCGAAGAAGACGGCTAAGATGCTGA A K L Y G P D V G Q P R R K T A K M L K 901 AGAGCTTCCTGAAGGCTGGTGGAGAAGTGATTGATTCAGTTACATGGCATCACTACTATT G E V 961 IGAATGGACGGACTGCTACCAGGGAAGATTTTCTAAACCCTGATGTATTGGACATTTTTA NGRTATREDFLNPDVLDIFI 1021 TTTCATCTGTGCAAAAGTTTTCCAGGTGGTTGAGAGCACCAGGCCTGGCAAGAAGGTCT V Q K V F Q V V E S T R P G K K V 1081 GGTTAGGAGAAACAAGCTCTGCATATGGAGGGGGGGGGCGCCCTTGCTATCCGACACCTTTG LGETSSAYGGGAPLLSDTPA 1141 CAGCTGGCTTTATGTGGCTGGATAAATTGGGCCTGTCAGCCCGAATGGGAATAGAAGTGG A G F M W L D K L G L S A R M G I E V 1201 TGATGAGGCAAGTATTCTTTGGAGCAGGAAACTACCATTTAGTGGATGAAAACTTCGATC M R Q V F F G A G N Y H L V D E N F D P 1261 CTTTACCTGATTATTGGCTATCTCTTCTGTTCAAGAAATTGGTGGGCACCAAGGTGTTAA LPDYWLSLLFKKLVGTKVLM 1321 TGGCAAGCGTGCAAGGTTCAAAGAGAAGGAAGCTTCGAGTATACCTTCATTGCACAAACA ASVQGSKRRKLRVYLHCTNT 1381 CTGACAATCCAAGGTATAAAGAAGGAGATTTAACTCTGTATGCCATAAACCTCCATAACG DNPRYKEGDLTLYAINLHNV T K Y L R L P Y P F S N K Q V D K Y L L 1501 TAAGACCTTTGGGACCTCATGGATTACTTTCCAAATCTGTCCAACTCAATGGTCTAACTC LGPHGLLSKS 1561 TARAGATGGTGGATGATCARACCTTGCCACCTTTRATGGARARACCTCTCCGGCCAGGAR

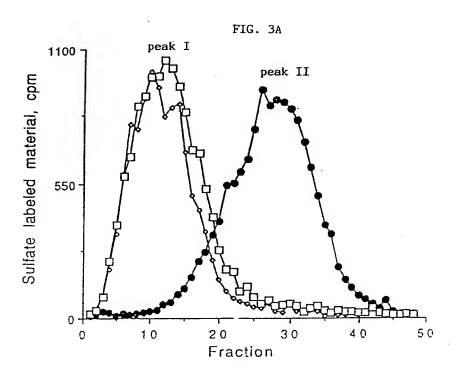
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A C 1

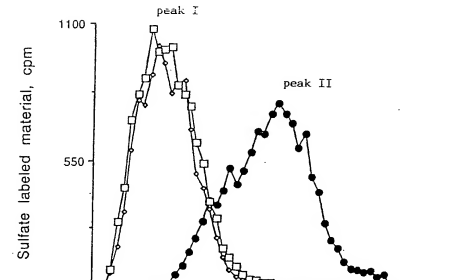
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1621 GTTCACTGGGCTTGCCAGCTTTCTCATATAGTTTTTTTGTGATAAGAAATGCCAAAGTTG
S L G L P A F S Y S F F V I R N A K V A

FIG. 2







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Fraction

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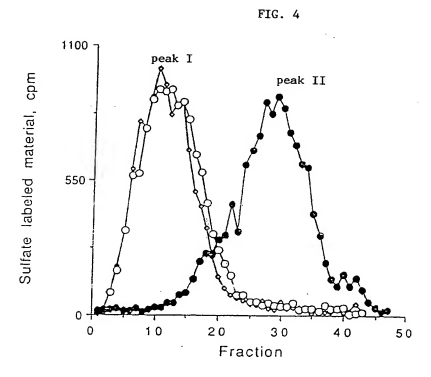
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FIG. 3B



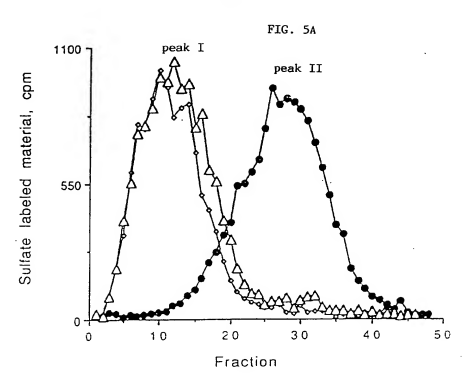
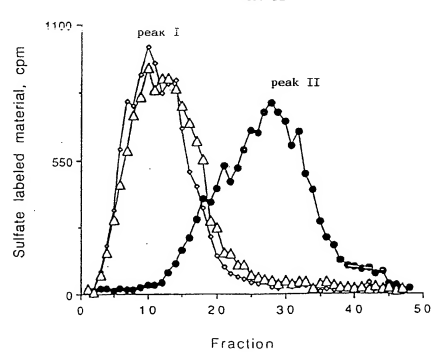
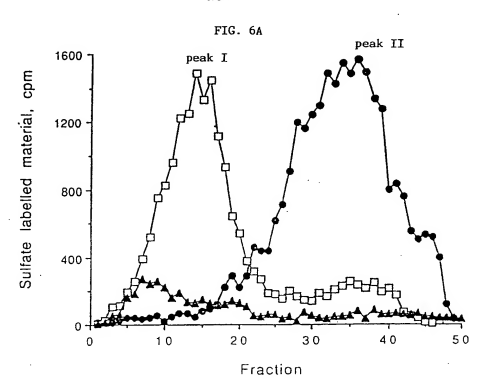
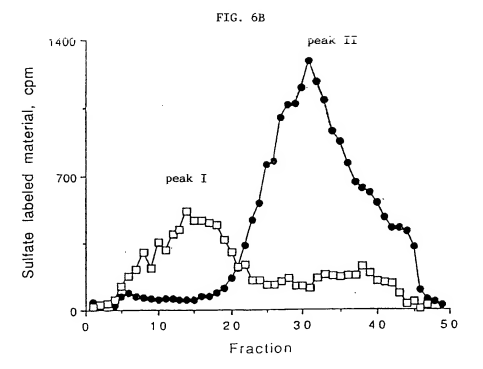
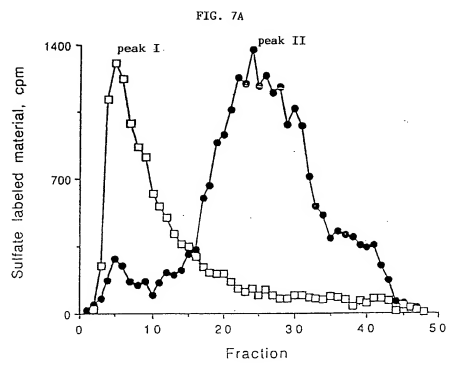


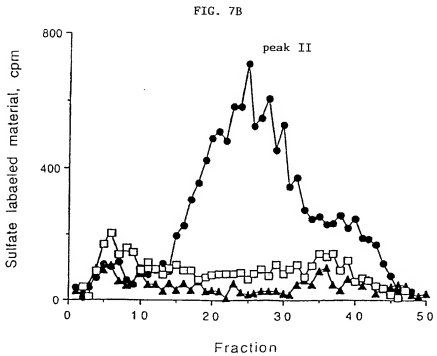
FIG. 5B



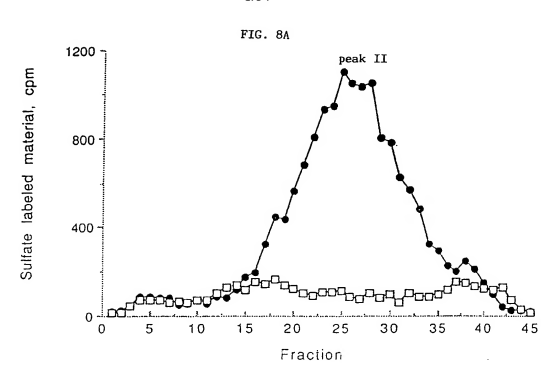












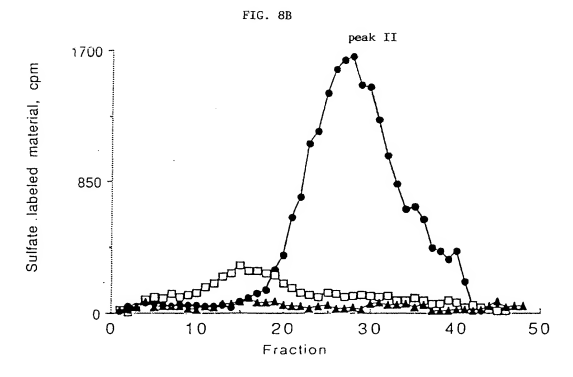


FIG. 9A

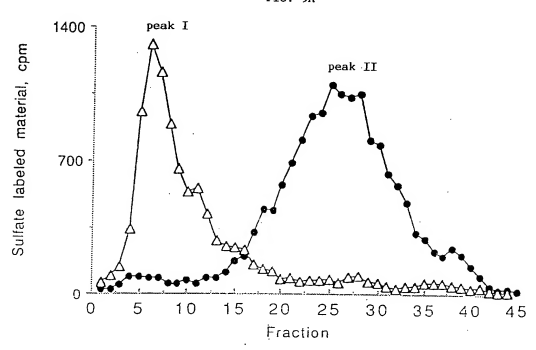
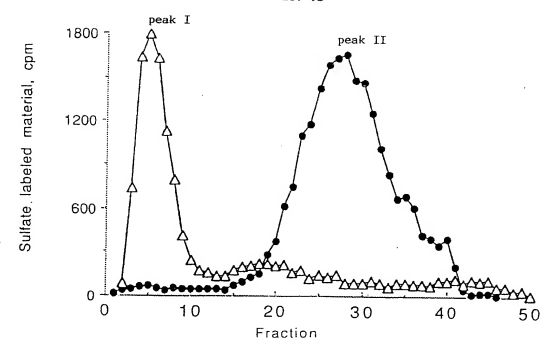
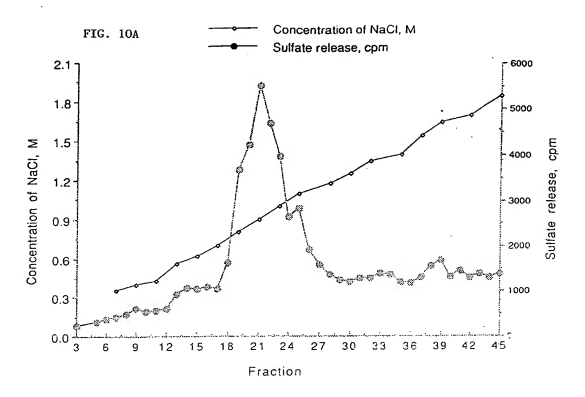
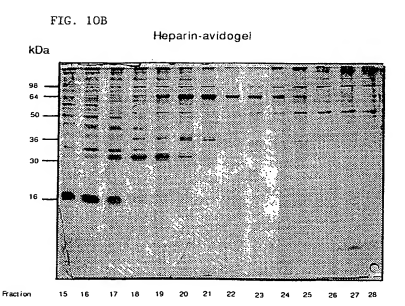
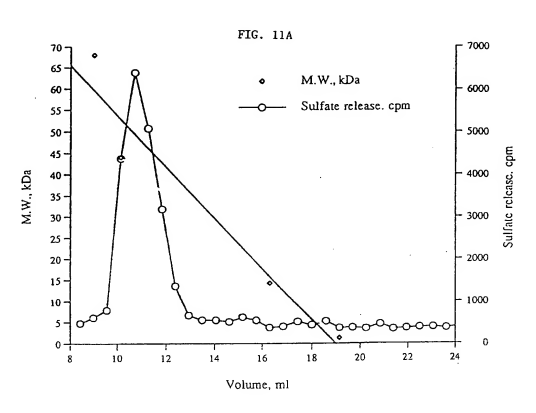


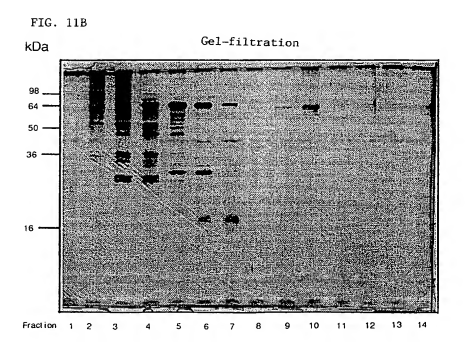
FIG. 9B











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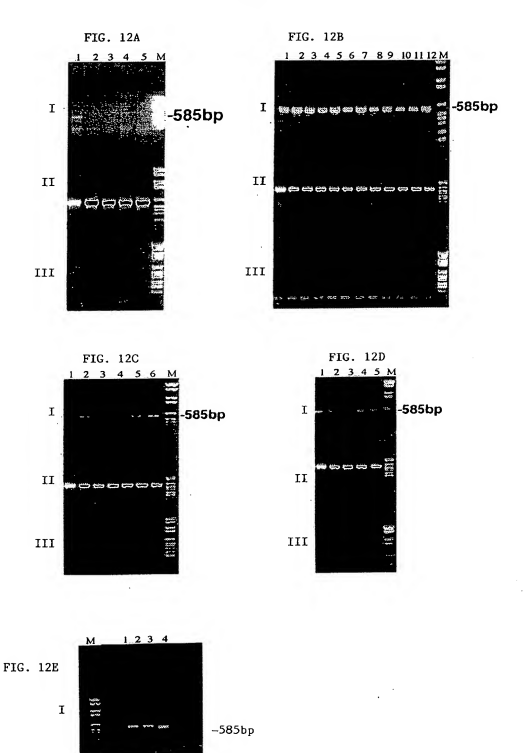
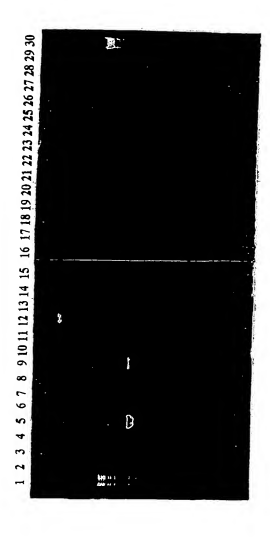


Fig 13

mouse	CTGGCAAGAAGGTCTGGTTGGGAGAGACGAGCTCAGCTTACGGTGGCGGT 5Q
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mouse	TOO
human	GCGCCCTTGCTATCCGACACCTTTGCAGCTGGCTTTATGTGGCTGGATAA 1165
mouse	ATTGGGCCTGTCAGCCCAGATGGGCATAGAAGTCGTGATGAGGCAGGTGT 150
human	ATTGGGCCTGTCAGCCCGAATGGGAATAGAAGTGGTGATGAGGCAAGTAT 1215
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human	TCTTTGGAGCAGGAAACTACCATTTAGTGGATGAAAACTTCGATCCTTTA 1265
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human	CCTGATTATTGGCTATCTCTTCTGTTCAAGAAATTGGTGGGCACCAAGGT 1315
mouse	GTTACTGTCAAGAGTGAAAGGCCCAGACAGGAGCAAACTCCGAGTGTATC 353
human	GTTAATGGCAAGCGTGCAAGGTTCAAAGAGAAGCAAGCTTCGAGTATACC 1365
mouse	TCCACTGCACTAACGTCTATCACCCACGATATCAGGAAGGA
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human	CTGTATGCCATAAACCTCCATAACGTCACCAAGTACTTGCGGTTACCCTA 1465
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human	TCCTTTTTCTAACAAGCAAGTGGATAAATACCTTCTAAGACCTTTGGGAC 1515
mouse	CGGATGGATTACTTTCCAAATCTGTCCAACTGAACGGTCAAATTCTGAAG 500
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FIG. 14



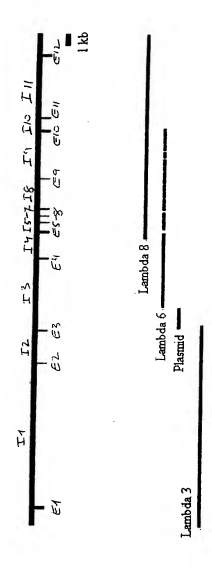


Figure 16

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aggaagcctggtccgggatgcccagcgctgctccccgggcgctcctcccc	2400
gggcgctcctcccaggcctcccgggcgcttggatcccggccatctccgc	2450
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aggggaaagcgagcaaggaagtaggagagccgggcaggcggggg	2550
ttggattgggagcagtgggagggatgcagaagaggagtgggaggga	2600
gggcgcagtgggggggtgaggaggcgtaacgggGCGGAGGAAAGGAGAA	2650
AAGGGCGCTGGGCGGGGGGAGGAAGTGCTAGAGCTCTCGACTCTCCG	2700
CTGCGCGGCAGCTGGCGGGGGGAGCAGCCAAGATGCTGCT	2750
M L L	
GCGCTCGAAGCCTGCGCTGCCGCCGCCGCTGATGCTGCTGCTCCTGGGGC	2800
	2000
R S K P A L P P P L M L L L G	2050
CGCTGGGTCCCCTCTCCCCTGGCGCCCTGCCCCGACCTGCGCAAGCACAG	2850

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P L G P L S P G A L P R P A Q A Q GACGTCGTGGACCTGGACTTCTTCACCCAGGAGCCGCTGCACCTGGTGAG 2900 D V V D L D F F T Q E P L H L V S CCCCTCGTTCCTGTCCGTCACCATTGACGCCAACCTGGCCACGGACCCGC 2950 PSFLSVTIDANLATDP ${\tt GGTTCCTCATCCTCGGGgtaagcgccagcctcctggtcctgtcccctt}$ 3000 RFLILLG tcctgtcctcctgacacctatgtctgccccgccagcggctctccttcttt 3050 tgcgcggaaacaacttcacaccggaacctccccgcctgtctctccccacc 3100 ccacttcccgcctctcattctccctctcccttactctcagacccca 31.50 aaccgctttttggggggtatcatttaaaaaatagatttaggggttacaag 3200 tgcagttctgttccatgggtatattgcattgtggtggcatctgggctctt 3250 3300 agtgtaactgtcacccgaatgttgtacattgtatctaataggtaatttct catccctcatccctctcccaccttcccaccttttggagtctccagtgtct 3350 actattccactaagtccatgtgtacacattgtttagcgcccactctaaat3400 gagcctttttgtttcattcattctgtaagtgttgaataggcaccacctaa 3450 ggtcaggtataagtggaaatttgaaaaagaaactgcccacttgccccagt 3500 3550 acttccctagccaagaggagggaaaccaggcaggtgcacctgaaggcctg tgagtgcttgatttgctgtgcagtgtaggacaagtaagattgtgcatagc 3600 $\verb"cttctgtatttaagactgtgttaggaagatttctcttttcttttct"$ 3650 ttttcttttttttttttttttttttaggcagatgaaaagggcgtca 3700 cagaacaggaataaaaatctaaatattcaataaatgagacctaggagact 3750 actgcagtgacttacaaagtcctaataaaaagatgtctctccaaaatggg 3800 gctgcaaaatgtggtgctgccttatcagctctaagttttttccttacctg 3850 agaaagaaggaacctgatgcaggttcagggctcctgccccatgaatgcag 3900 gctgactccaagatggggagctacagggacaatcccaggtcttctaggcc 3950 tcttatttaggccctgggagcctccagagatggccacatcttgaccagcc 4000 cagatagagggaaagatcaccattatctcacctctgtgtcaaatacctag 4050 atgctgtcctcctgagcccacactatagttgccagcgctaatttaatgg 4100 gtagtgtactggttaagagatggacagaccatcctggcttgactctcagc 4150 tctggcaaagatgagtgacttggtttttccatatctcttggccacaccaa 4200 ccttgatttcttcagctgtagaatggaatttctcaagcttgcctcaagga 4250 ttattgcccgaggatttgatgatatggtaagagcttctcagtgtttgacc 4300 4350 4400 tgagcatttggtagccattcaccggttttctgtttcttttggatcatagtt 4450 aacctctccttttccttctggcactacaattttctggtggggaagaatcc ttactttstgsssttssscttaaggataggaagstgatastaggsagsaa 4500 ctagttgggggataggaagattgttccagagaaatgctgaaccatagggc 4550 4600 gggcggttactgaacatgggtatgaagtagatgtccatttactgaaatgt 4650 gaggacctgaggcctcttctattgctgtagccagcatattccccaacctc 4700 tccccaagaaaggacagatgggggttccccctggagtaacaggtccaaa 4750 agaaaaaacatacagtgggacttccaggatctgggcctgatcacccagca 4800 gtcaagctccccgcaattgactaacacccccctaacacgtagaaattcca 4850 atctgcaatttagtgaggatgatacctttattcttcttaaatacatctct 4900 4950 tcatttcccagagcacccttttttcccctctctgcacctttttgttaaa gactggagtataatgaaataccaagagagcataacatgtgatacataaaa 5000 5050 ctttttttctggtttacaaaacagttcattcttgtccatacgtgcttctc tccaaggctggctgtctgttccagcccgcttcgcttggagaggccat 5100 $\verb|ctgccatacctgctccccagacgcatcgacaagcacacccagagtgttat|\\$ 5150 $\verb|ctgctaagacctaaaagagggaggaaccccctctcctcatctaagaccta|\\$ 5200 gcttctaaattagagtgtgagggtccatctccccaggagggcacagggc 5250 $\verb|cca| a a cagc c cagc catct c a gaaga c a a cacta a g cttt g t a g g g t c c$ 5300 acagtagaggagagtaagacgcctgttgtttaatttattacagttcctca 5350 5400 aaagtgaagatgtgtggggggggggatggcaagagctgagcagacgaaagctg 5450 aaggaataaggaaagaggaggacacaaacagctgacacttcctcagtt 5500 $\verb"cttgtcatttgcctggccctgttctaagcaccttctaggtattaatccat"$ ${\tt ttagtcttggctacaacactgtgagtaactagttttgtcaccccatttt}$ 5550 aaaaatgaagaaagtgaggctcagggaggttaagtaacttggccacagtt 5600 tgaaactagactctgatcacatgagataatagtgcccataaaaagggaaa 5650 gcagattatatttttaaaggaaagagtaggatatggtagaaaaagat 5700

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Fig. 16 (continued) aggagtccctctgaatgtttccatagcatttttaaagaattgcctattta 11850 cttgttcgtatctatcactaaactacaaattgtatgagaacagccactat 11900 ctctgcctggttcaccattcatctccagcaactagcataatgcctggcag 11950 agtcagcctgcaacaatatttgttgaataaattaacagatggctttatc 12000 tccttaagtaaatcttgcttttttcacctattaaaacagacgcacaggcc 12050 aggtgtggtggcccatgcctgtaatcccagcactttggcaggctgaggtg 12100 ggcggatcacctgaggtcaggagttcaagaccagcctggccaacatggtg 12150 aaaccccatctctaataaaaatacaaaaattagctgggcatggtggtggg 12200 tgcgtatagtcccagctactagggaggctgaggcaagagaatcgcttgaa 12250 cccaggaggcagaggtggcagtgagccgagatcatgccactgtactccag 12300 12350 cacacacacacacacacacacacagttgtataatttaaaata 12400 taacgtgcttgttatggaacacttgtaaaatacaggaaagtaatgaaaaa 12450 gtctaccatctagctcaccacataatgaccattgctatcatcctggcata 12500 attetetetgtatataaatatattettttattgttaaaattacacta 12550 tgagtactatttattttattttactgtggcaaaatgcgcaaaacataaaat 12600 cttgccattttaaggtatgcagtttggtgcattcaccacactcacattgt 12650 tgtgcaaatatcaccactatctatctcagaacttcttcgtcttcccaaac 12700 tgaaactctgtacccattaaacaatagtgcatcctctgttttcccctccc 12750 tacaatttattttatttgggtttgtaccaaactgaaaatagctgcttct12800 tccttacttagttcagattagcatttccatttatttagccgtggttttga 12850 ggatgccatgacagatgccatccttcctagagctctttggggctgtcagg 12900 ${\tt tatttcagtcagggtgaattcgggttgataacattttaaaatctcacttt}$ 12950 attctgaggttcctagtgtcagagcccaccgtatttttagggactcccaa 13000 gttacaaacaaaatatggtgaggaggaatcactgaagttttaacacaag 13050 agacttacattttgttcaatttctatcttttagtttatttcctaagcata 13100 13150 tgagcacatcttaaaactttaaattttagatcagatctttaattcctagg 13200 13250 atcccaacactttgggagggtgaagtgggcgaattgctagagcccaggag 13300 gtggaggctgcaatggcctgagatcacgccatcgtactccagcctggatg 13350 13400 gaagaagtattggcaatcagtgctccaggaataatttcctgacttgaaat 13450 aaacctacatgtagacaaactaattaggccattccaagagttgctagcat 13500 ${\tt tggtttaatatgttttcagagcattccaggaagcagtgtggccagcattg}$ 13550 catgtttgatacttcagaaatgtatgacaggtgtttctcttacccaggtc 13600 ttctgttttcttagttttgctcatgtaaatatttatgaacatcctcatct 13650 ${\tt ttttgagggaagggattatagatcattctaattccattttctagcatttg}$ 13700 gtaccattctaagcacatgataggcacccatttggagcatttttggcttg 13750 acagaatatgcatttagaattgttcaaattagaggtgtcagtgatgggaa 13800 ttagaatactatataattctaagtcatttgacttaaatacaaaagaatga 13850 ttttccttggtggggaatggtgaagggaggcaggagttaagaagaggaga 13900 agagatcctaagtcatttataaacttctctggaaagacaggtgtgtgaag 13950 14000 $\verb|ttaaatagactttatttttagagcagttttaggttcacagcaaaattga|\\$ 14050 14100 14150 gaacctacactgacacatcattatcacccaaagtccatagttcacggcag 14200 ggttcactgtcggtgtacattctatgggtttgagcaaatgtataatgaca 14250 tgtatccaccattatagtaacatacagagtattttcagtgccctgcaaat 14300 ccctgttctccacctattcatccctcctctctgcatttccacccccag 14350 cccctggtaaccgctgatctttttactgtcccatagtttcggacgatcta 14400 ${\tt tttttcagacagacacagagctgtctttcccttagtttctatcat}$ 14450 14500 14550 tggctcatgcctgtaatctcagcattttgggaggctgtggcagaagcatc 14600 acttgaagccagaagtttgagaccagcctgggcaacatagcaagacccca 14650 14700 cacacacacacaaaacaagctcttgccagaattagagctacaaattg ccctcaggttcctagaagatcagtccttcaattagattcagattgagatg 14800 cttcctcttttaaacaatgattccctttctatcatgcccaataagaaaac 14850

	Fig.	16(contin	ued)		21/3	34		
222	-							
220		ttaaacaat	actgcc	tgtaato	ccagcta	cccagg	aggca	g 14900
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caaa	acaat	cactccago gtgatttco	tectet	aacagag	caagatt	ctgtct	caaaaa	15000
aata	agtcca	accaggaaa	gaagga	antaana	atattta	gaaatg	ttaaga	15050
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GCTT	CGTACC	CTTGGCCAG	AGGCTT	GTCTCCT	GCGTACC	アピタピになっ	המהכיכהים ב זי	15200
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GCAC	CAAGAC	AGACTTCC	TAATTT	CGATCC	CAAGAAG	GAATCA	r ማርርብ	15250
GI	' к 1	'DF	LI	FDP	KK	E S	т г	-
GAAG	AGAGAA	GTTACTGG	CAATCT	CAAGTCA	ACCAGGa	tgaaaat	ttttta	15300
E	E R	SYW	Q S	Q V	N Q			
aaga	ttcact	ctatattt	taattaa	cgtcag	tccgtca	tqaqaat	acttt	15350
gaga	aaactg	ttatttct	cacacct	aacaat	taatgaga	attaact	tecte	. 15400
tccc	ctcatc	tgacctgt	ggaggaa	tctgaa	caagagga	aggagge	agtgg	15450
gcag	gtttcc	ttatcatga	atgtttg	rtcatgt	tcagtgto	gaggcct	cacaa	15500
aaaa	aaaaaa	aaaaaaaa	aaggcgt	cctggai	tataacto	gagaget	catto	15550
taca	gtaaat	attaataaa	acagtg	rattgtag	gctgaag	gatagaa	ctgct	15600
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caaa	gacaca	atgatagat	tgaagg	atattta	ttctaaa	atataga	atatg	15700
ggtg	aacgag	atctgtgga	icttctg	ggctcca	acgttag	gattctg	atttt	15750
agca	agcttg	tcaggggat	tctgat	attgaaa	iggctgtg	gccttc	acctg	15800
agaa	acctge	cctaggggg	ccatga	aaatttg	tcctgtc	tttcag	aagtg	15850
gggg	rayaca	tcaaatgga	lagttaa	accgtat	cttaaca	attact	aggat	15900
gggcg	rateac	actcacacc	cytaat	cccaaca	ctttggg	aggetg	aggca	15950
agage	ttatal	ttgagccca tctatttt	taataa	tttaaac	agcccgg	gcaaca	tagag	16000
tatto	tataca	accactgaa	ttataa	taatoto	tatataa	tatatt	gaaaa	16050
attat	gaggaa	atatttgat	tatttc	atatatt	atatett	ttcctt.	ctatt	16100 16150
tattt	tatcca	agttatgaa	gtattt	agaacaa	ttcatca	gtaatt	naaac	16200
taaat	tgacac	gaatagtaa	tcagaga	aaatag	aaaaaga	cagato	ggggc	16250
tcttt	gaatad	caggitgg	agttgti	ttatggg	tttattt	tttatt	ttaaa	16300
ggcgt	ttttt	agacagag	téccaet	ctgttg	cccaggc	tagagt	gcagt	16350
ggcac	aagcat	ggcccact	gcatect	tgacct	cttgggc	tcaage	aatct	16400
tccca	ccttag	cctcctga	gtagcto	ggacca	caggtgc	atgtcad	ccaca	16450
cccag	ctaatt	ttttttatt:	ttttgta	igagaca	gtottto	tatgtta	atcca	16500
ggctg	atctca	aactcctg	cactcaa	gtgatc	ccctgc	cttggc	gtccc	16550
aaagt	attggg	attatagg	catage	caccaca	cccaacc	tagttt	ctatt	16600
tagac	ttggcc	ctttccca	ccagtca	tttgtg	ccaaaa	gatetea	taaa	16650
Lgtag	acagga	aactgtcci	tttgctc	atcagt	ttcttca	atcctgt	gtct	16700
agggg	gatggt	cggtgggg	gaaactg	gggttal	gcaagt	tcctctc	aaac	16750
tatac	cigiga	gcccaggga	atggatg	aggcaco	cagccgc	cagcgag	tcag	16800
gccag	cacccc	Ccagaaagg	jaagtea	teageca	igtcagco	ggccct	ggca	16850
acagg	ataata	gcaacccto tggatttt	ctttt	+c++++	:aayaaa:	-ggtete	cctg	16900
ateta	acteta	tcgcccago	ctagaa	tacaata	.cccaaaat	ctyaga	cagg	16950
tgcag	cctcta	cctcccag	ctcaaq	gcaticct	cccacct	caatet	ccac	17000 17050
agtag	ctagga	ccacaggca	cacacc	accacac	ccaacta	aattt	cate	17100
ttttt	agtaga	ggcagggtt	ttacta	tattato	caggeta	atctca	aact	17150
cctga	gctcaa	gctatccat	ctgcct	tagecte	ccaaaga	gctgga	atta	17200
caagc	gtgagc	cactgtgcc	tgacca	gggtgga	ttttttc	aagtgc	acat	17250
gttgt	ggtccc	agaagctct	gatggt	accaaat	tccaago	gaaaaa	aagt	17300
caatg	gttccca	acccatcct	acctcc	catgatg	gcaagag	gaaatc	acca	17350
cactg	cagata	cagtccatg	taaaac	aaattgo	tatqqat	tttqaa	agtg	17400
aaccti	caagaga	actgcact	atgttt	tcttcat	tagagtt	ctctaa	taat	17450
ttcca	gctttti	tttttttt	tttttt	agacagt	gtctcgc	tttqtc	accc	17500
agtgto	cacccag	gctggagt	gcagtga	acgtgat	ctcggct	cactac	aacc	17550
teege	tcgtg	ggttgaagt	gattct	cctgcct	cagcctc	ctgagt	agct	17600
gtattt	cagtag	gagacgagg	tttcac	catttgg	ccagget	ggtctc	gaac	17650
rectga	ccccaa	gtgattcg	cccatc	ccagcct	cccaaag	tgctgg	gatt	17700

Fig. 16 (continued)

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ataaatagctaactaaataaatatatgagatttcagtctgctcactgtga	18000
aaatagaccttctaaatgatctcttccacttgcagATATTTGCAAATATG	18050
D I C K Y	10030
GATCCATCCCTCGATGTGGAGGAGAAGTTACGGTTGGAATGGCCCTAC	10100
 	18100
G S I P P D V E E K L R L E W P Y CAGGAGAATTGCTACTCCGAGAACACTACCAGAAAAAGTTCAAGAACAG	10150
	18150
CACCTACTCAAgtaagaaatgaaaggcaccctagagatgttccagcccca	18200
T Y S	
aagatatttgaataggttggactcgggcaccaatctagcaagtcctacgg	18250
aagttgtataaagctgaaaatactgaagcatttcccaaatgggaaatcct	18300
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ettgaatctgggaggcagaggttgtggtgagccgagattgcgccaccgca	20550

20600 aaaacaaacaaacaaacaacaacaacaaaaaaaacgggtatcccagaa 20650 gatacaggtaagttttctaacacaggtcctcttgtatggtgcgttccact 20700 taagtagaagatgacaaaaacatttgtcatgagaatatagactcacattt 20750 taaacctgtttgagcaggaaaaggaagcaatgttacagatgtaattctgg qtqtqactgcagaaaggatgactcccttattaaagtagtcatcctqaqtq 20850 agctaactctttgtacttcctcttctcctctgttcccctcatcacccca 20900 20950 ttcttccgttgcctacacccaggcccacattggatgctgacatagactta catggtacagtccaagggaaagatctgccatttttttcaatgtgtcatct 21000 tggttatcttcattccaaggatctctccactctttatacagtaagagatg . 21050 agagtctggaaaggattgggaataagataatgaattgtaagttttaaatt 21100 $\tt gttcttcgtattttggggaaggagtaggctaggtggtccttctgttttt$ 21150 ttttqtttttttttaaaqtaqatqtqqccaqacqtqqtqqctcacqcc 21200 tgtaatcccagcactttgagaggctgaggcaggtggatcacttgatgtca 21250 ggagttcaagaccagcctggccaacacagtgaaaccccgtctttactaaa 21300 aatacaaaaactagccgggcttggtggcgtccacctgtagtcccagctac 21350 tgcagaggtggaggcaggagatcacttgaacccgggaggtggaggttgc 21400 agtgagccaagatcatgccattgtactccagcctgggcgacagaacaata 21450 ctctgtctcaaaaaaaaagaaaaagaaagaaagaatggatttga 21500 actcagtcgtcaatagcctctattccaggagatgttacagttgattatgt 21550 tatagggggtgtataatagaatttcgagctatgtaaattccaagtgcatt 21600 tggaagaatgaagaatggaggaagggtaaagtatgagtgcaagcattcc 21650 aggttttttgaaaatgctataatctttgttcagggctagtacaaagtgct 21700 21750 atttagctgtaagggttttttgtgatttacagacagttttcacatgtgtc atttcaaccttggttttatggcgaaggcatgtgatggtgcttgtcccagg 21800 actttagatccatatctgaggttcctgtcgggcaaagatattacccctga 21850 21900 tcatattatagtctataagtgggagagttgtgcctggagctcaagtctta 21950 tgatttctgatccagggcacttcctacaacatgattttgcaatataaaag cctataatgtgtgactaaagcaggtcactcaccccttgtaacagactcta 22000 gtaatggtactgccaccaaacggctgcgtgatattgggcaaagacttacc 22050 ttatttgaatctcagtttcctcctagaaaaatgagggtggaggttaagca 22100 taggctgatgatcctaaagcctccatactgccctaaactgtggctctaag 22150 atccagtagaatgctgggtcacaggactctagggagcttttcaaacccaa 22200 atgtctgtcattccttgatggtaggcagcagtttatggaagtgggcgaca 22250 22300 caqcaaatatcaaaatacctaaaqcaqcttqcaaqaqttqtttctqccta 22350 agagtcttgctctgttacccaggctgcagtgcagtggcacaatctcggct 22400 cactgcaacctccacctcccgggtttgagcaattctgtctcagcctccca 22450 22500 agtagctgggactacaggtgcatgccactgcacccagctaatttttgtat ttttagtagagacggggtttcaccatattgggcaggctggtctcgaactc 22550 ttgacctcaggtgatccacctgcctcagcctcccaaagtgctgggattac 22600 22650 aggcatgagccactgcacccagcttaaatagctaatatttaatattattc 22700 tataqttattcaaqtaattcaqqccaaaqacttaqaaacaaaacaaaaq 22750 cttttttaactacaagagttcaggaatgaattactctttaacaaacgact 22850 atagatatacatgaaaattggaaggacttattatgcatatgataatcaat ttaaagacaacacttaaaattatattgttgccactctcaaaaagtggtaa 22900 tagaacagctaatggtttaaaaagcagagtacagaagttcccaaacttat 22950 ggcaccttaatatcgcagaaaactttttaaagcatgcctaggccacaaaa 23000 aatacctgtattttgattattaaattgtaaggtctacacaacctaatagt 23050 aataggtccaatagtaatgctgtccaatagatgttgatgtttttttcctt 23100 gcaaacttaaaagatcctacagtgcctctgtaaatagcactgcctggtta 23150 23200 qaqttqaatttcaqataaataatttttttcatqttaattatttttcttt 23250 gggtctcattctgttgcccaggctgctgtgcaatggcatgatcatggctc 23300 actgcagccttgacctccctgggctcaggtgatcctcccacctcagcctc 23350 23400 ccaagtagctagctgggactacaggtgcttaccatcatgcccggctaatt tttgtgttttttgtagagatgtggttttgccatgttgcccaggctggtct 23450 tgaactcctgggctcaagtgatccgcccgcctcggcctcccaaagtgcta ggatgacaggcatgagccactgcacctggcccctgggcgaagtatttctt 23550 aatggttacataggacatacactaaacattatttattgtctatatgaagt 23600

25600

25700

25750

25800

25850

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25950

26000

26050

26100

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26350

26400

26450

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Fig. 16 (continued)

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tgtaatttggaaaatgttttgttgcttttaaatactgctgtatgtttgct	26550
tttaaatacaacatttctcgatatatattttgagaattgctgtctttcag	26600
AACCTAACAGTTTCCTTAAGAAGGCTGATATTTTCATCAATGGGTCGCAG	26650
E P N S F L K K A D I F I N G S Q	
TTAGGAGAAGATTTATTCAATTGCATAAACTTCTAAGAAAGTCCACCTT	26700
LGEDFIQLHKLLRKSTF	
CAAAAATGCAAAACTCTATGGTCCTGATGTTGGTCAGCCTCGAAGAAAGA	26750
K N A K L Y G P D V G Q P R R K	
CGGCTAAGATGCTGAAGAGgtaggaactagaggatgcagaatcactttac	26800
T A K M L K S	26050
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Y	28000
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Y L N G R T A T R E D F I N P D V	20050
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V V E	
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S T R P G K K V W L G E T S S A Y	
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ggatgaattgtttcagaaattttggcctttaattatggcccataaatat	28900

Fig. 16 (continued) gtcaagtagtccttactctaaagaagtacactgtaaaagaatgcatatag 28950 ccggatatggtagttccctgtaatcccaatactttgggaggccaaqqtqq 29000 gaggattgcttgagcccaggagtttgaggctgcagtgagttatgatggtg 29050 29100 ctctgtcacccagactggagggcagtggcacgatctcacctcactgcaac 29150 ctctgcctcccggattgaagcgattctcctgcctcagcgtcctgagtagc 29200 tgggactacaggagtatcaccgcactgggctaatttttgtatttttagta 29250 gagacggggttttgacatgttgcccaggctggtctgaaacccatgagctc 29300 aagtgatctqcctacctcaqccttccaaaatqctqqqattacqqacatqa 29350 29400 29450 ctataattcatagattcccaagaagtttagagcctaaagtatgaggtccc 29500 accagaggggctatcattaaatttaaagatttgttaaatcatctcattgt 29550 ccaacaccacaaacttgattgctttaaaatactggtttagttacatttag 29600 taactctattagtgcttttaatctatactgctatatcctcacattgagat . 29650 29700 ttataagcctagaatacatcacaaatcctttatgcccatggaagcaagag 29750 gaataaagaatggagatgtttgttttgccattaactaaagatctggggtg 29800 tcggggagaaggggatagagaaggagaagtgggaagaggtgtccataat 29850 agcttaggtgcaattctgcttattttacattttacccccgctgactgcca 29900 ctttttcttcagccctcacacattgtttgtgcagggacctcataggacca 29950 ggaattgtctatagaggtgggaatttgtctcaccctgaaagggatacctc 30000 tagcatggtaatagtcttctaggatttgttatcatatggaaagatgtaaa 30050 gggagggattctgctgctgctgctgctgcatgcagttgccatttcat 3 100 ttaaatgacttatttataattgatgacacttttctggcttcctgttaatt 30150 cctccctcaaagatcaataaaccagaaccaggcatggtggcatgcacttg 30200 30250 ccaattatcaagacaggggaattgcaaaggagaaagagtaatttatgcag 30300 agccagctgtgcaggagaccagagttttattattactcaaatcagtctcc 30350 ccgaacattcgaggatcagagcttttaaggataatttggccggtagggc 30400 ttaggaagtggagagtgctggttggtcaggttggagatggaatcacaggg 30450 agtggaagtgaggttttcttgctgtcttctgttcctggatgggatggcag 30500 aactggttgggccagattaccggtctgggtggtctcaaatgatccaccca 30550 gttcagggtctgcaagatatctcaagcactgatcttaggttttacaacag 30600 tgatgttatccccaggaacaatttggggaggttcagactcttggagccag 30650 aggctgcattatccctaaaccgtaatctctaatgttgtagctaatttgtt 30700 agtoctgcaaaggtagacttgtccccaggcaagaagggggtcttttcaga 30750 aaagggctattatcatttttgtttcagagtcaaaccatgaactgaatttc 30800 ttcccaaagttagttcagcctacacccaggaatgaagaaggacagcttaa 30850 aggttagaagcaagatggagtcaatgaggtctgatctctttcactgtcat 30900 aatttcctcagttataatttttgcaaaggcggtttcagtcccagctactt 30950 gggaggctgagacaggaggattaatggagcccaggagtttgaggttgcag 31000 agagctatgatcacgccactgcactccagcctgggtgacagagtgagacc 31050 31100 aagatggtgtgcaattagaattgagcgattttgtttccaaacctcaagaa 31150 agcttggtcttgctctgtcccagGTGGCTGGATAAATTGGGCCTGTCAGC 31200 WLDKLGLSA CCGAATGGGAATAGAAGTGGTGATGAGGCAAGTATTCTTTGGAGCAGGAA 31250 RMGIEVVMRQVFFGAG ACTACCATTTAGTGGATGAAAACTTCGATCCTTTACCTgtaagtgaccat 31300 NYHLVDENFDPLP ${\tt tattttcctaattctagtggagtagattaaagtcaactcaggacctctgg}$ 31350 tgttaacctcctatgaacagtcagtcctctcagtaactagccaaatcatg 31400 agatgatgaattagaaggagccttagatagcatccaatctaacatttttt 31450 tgtgtgtttgaagagaagaaatcaagagctaggaataactttttaaaggt 31500 aagccatttgcagtatagtgtggattttgtttaaaagggggataatttgaa 31550 attttatgactcattatacaagacaaaataagttggattttcaaatgttt 51600 tacaaagtaaatcaaagttataattgcctacagtacgcaaagcttcaaaa 31650 cattttttatgttatgaaattgtaatttatttaaccttaaaatgagccag 31700 taccatgtgtttgcttaaaaatctcatgctaagaatttactatgttgtta 31750 31800

Fig. 16 (continued)

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K. E. G., D. L. T. L. Y. A. I. N. L., H. N., V. T. K.	
TACTTGCGGTTACCCTATCCTTTTTCTAACAAGCAAGTGGATAAATACCT	36500
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TCTAAGACCTTTGGGACCTCATGGATTACTTTCCAAgtaagtaattttcc	36550
L. B. P. L. G. P H G L L S K	
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agtttggacagggagcaaaagacaaagtcaactatatcaagttctaataa	36650
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43250

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30/34 Fig. 16 (continued)

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Fig. 16 (continued)

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ttccattcaatttcattcaatttaagtatatttgtaaggagctaaagct	tg 44650
aaaattaaattttagatctttcaatactcttaaattttatatgtaagtg	gg 44700
tttttatattttcacatttgaaataaagtaatttttataaccttgatai	tt 44750
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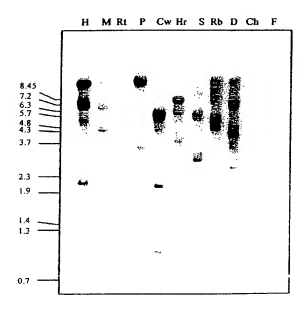
32/34

Figure 17

.	MINGON				50
human					LDFFTQEPLH
mouse rat					LEFYTKRPLR
Iac		LDLLWLWGI	LINALIQUIPA	GIAPINDOVD	LEFYTKRLFQ
					100
human	LVSPSFLSVI	IDANLATOPE	FLILLGSPKL	RTLARGLSPA	
mouse		IDASLATOPE			
rat		DASLATOPE			
					150
human		TFEERSYWOS			
mouse		TSEERSYWKS			
rat	FILEDPNNEP	TSEERSYWOS	QUNNDICGSD	RVSADVL~~~	~~~~~~
					200
human	LLREHYOKKF	KNSTYSRSSV	DVLYTFANCS	GLDLIFGLNA	
mouse		KNSTYSRSSV			
rat		~~~~~~~			
_					250
human		CSSKGYNISW			
mouse	_	CSSKGYNISW			
rat	~~~~~~	~~~~~~~	~~~~~~~	~~~~~~	~~~~~~~
					300
human	KLLRKSTFKN	AKLYGPDVGQ	PRRKTAKMIK	SELKAGGEVT	
mouse		AKLYGPDIGQ			
rat		~~~~~~~			
					350
human		LNPDVLDIFI			
mouse		LSSDALDTFI	_		
rat	~~~~~~	~~~~~~	~~~~~~~	~~~~~~	~~~~~~
					400
human	GAPLLSOTEA	AGFMWLDKLG	I.SAPMCTEVV	MPOVERGACN	
mouse		AGFMWLDKLG			
rat		~~~~~~	_	-	
					450
human	LPDYWLSLLF	KKLVGTKVLM	ASVQGSKRRK	LRVYLHCTNT	DNPRYKEGDL
mouse		KKLVGPRVLL	SR V K G PD R SK	LRVYLHCTNV	YH PRYQEGDL
rat	~~~~~~~	~~~~~~	~~~~~~	~~~~~~	~~~~~~
					500
human	TT.VATNI:UNT	TKYLRLPYPF	CMECADEUT	DDI CDUCT I C	500 KSVOLNGLITI
mouse		TKHLKVPPPL	_		_
rat		~~~~~~~~~			
.=		-	-		
					543
human	KMVDDQTLPP	LMEKPLRPGS	SLGLPAFSYS	FFVIRNAKVA	ACI~
mouse	_	LTEKPLPAGS			
rat	KMVDEQTXPA	LTEKPLPAGS	SL SV PAFSYG	FFVIRNAKIA	ACI~

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Figure 18



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Figure 19

	IMLLRSKPAL	PPPLMLLLL	GPLGPLSPG	ALPRPAQA(ODVVDLDFF"	PQEPLHLVS	PSFLSVT	1 6
PHD	1	EEEEE			нин	EEEE	EEE	I
PHD	(IDANLATDE	RFLILLGSPI EEEEE	KLRTLARGLS	SPAYLRFGO HHHHE		PKKESTFE		
PHD	LEEE	EEEEE	нинин	нинк	EEEEE		нинин	
PHD	I QVNQDICKY		CLRLEWPYQE IHH HHHH					
	,	10000				CCCCCCCC	EEEE (
P.HD .	GLDLI FGLN				niswelgné Eeeee		•	
	QLGEDYIQLE				КМLКSFLKA НИНИНИНИ			300
							•	
	NGRTATREDF				_	SAYGGGAPL	LSDTFA	360
PHD.	i	нинин	нининеее	EEEE	EEEEEE	H	मम्म्ममा ।	
	AGFMWLDKLG						•	420
PHD	нининин	нин ини	нининин	EEEEE	ннни	нининин	EEEEE	
	ASVQGSKRRK						•	480
PHD	EEE E	EEEEEEE		EEEEEE	EEEE	Е ни	нинини	
	RPLGPHGLLS			PLMEKPLR	PGSSLGLPA		•	540
PHD	nn EE	eeeee ef	EEEE			EEEEEE	ee ee	
DIID	IACII							543
PHD	1 1							

1 SEQUENCE LISTING (1) GENERAL INFORMATION: (i) APPLICANT: Iris Pecker, Israel Vlodavsky and Elena Feinstein (ii)TITLE OF INVENTION: POLYNUCLEOTIDE ENCODING A POLYPEPTIDE HAVING HEPARANASE ACTIVITY AND EXPRESSION OF SAME IN GENETICALLY MODIFIED CELLS (iii) NUMBER OF SEQUENCES: (iv) CORRESPONDENCE ADDRESS: ADDRESSEE: (A) Mark M. Friedman c/o Anthony Castorina (B) STREET: 2001 Jefferson Davis Highway, Suite 207 (C) CITY: Arlington (D) STATE: Virginia (E) COUNTRY: United States of America (F) ZIP: 22202 (v) COMPUTER READABLE FORM: MEDIUM TYPE: (A) 1.44 megabyte, 3.5" microdisk (B) COMPUTER: Twinhead* Slimnote-890TX (C) OPERATING SYSTEM: MS DOS version 6.2. Windows version 3.11 (D) SOFTWARE: Word for Windows version 2.0 converted to an ASCI file (vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: (B) FILING DATE: (C) CLASSIFICATION: (vii) PRIOR APPLICATION DATA: (A) APPLICATION NUMBER: 08/922,170 FILING DATE: (B) 2 SEP 1997 APPLICATION NUMBER: 09/109,386 (A) (B) FILING DATE: 10 JUL 1998 (A) APPLICATION NUMBER: PCT/US98/17954 (B) FILING DATE: 31 AUG 1998 (A) APPLICATION NUMBER: 09/258,892 (B) FILING DATE: 1 MAR 1999 (viii) ATTORNEY/AGENT INFORMATION: (A) NAME: Friedmam, Mark M. REGISTRATION NUMBER: 33.883 (C) REFERENCE/DOCKET NUMBER: 910/62 TELECOMMUNICATION INFORMATION: (ix) (A) TELEPHONE: 972-3-5625553 (B) TELEFAX: 972-3-5625554 TELEX. (C) (2) INFORMATION FOR SEQ ID NO:1: SEQUENCE CHARACTERISTICS: (A) LENGTH: (B) TYPE: nucleic acid (C) STRANDEDNESS: single TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1: CCATCCTAAT ACGACTCACT ATAGGGC 27 (2) INFORMATION FOR SEQ ID NO:2: SEQUENCE CHARACTERISTICS:

(A)

(B)

(C)

LENGTH:

TYPE:

24

STRANDEDNESS: single

nucleic acid

				2
		(D)	TOPOLOGY:	linear
	(xi)	SEQU	ENCE DESCRIPTION	: SEQ ID NO:2
		GTAG	TGATGC CATGTAACT	G AATC 24
(2)	INFO	RMATION	FOR SEQ ID NO:3	:
	(i)	SEQU	ENCE CHARACTERIS	TICS:
		(A)	LENGTH:	23
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUI	ENCE DESCRIPTION	: SEQ ID NO:3
		ACTC	ACTATA GGGCTCGAG	C GGC 23
(2)	INFO	RMATION	FOR SEQ ID NO:4:	
	(i)	SEQUE	NCE CHARACTERIS	TICS:
		(A)	LENGTH:	22
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUE	NCE DESCRIPTION	SEQ ID NO:4:
		GCATO	TTAGC CGTCTTTCT	r CG 22
(2)	INFOR	MATION E	FOR SEQ ID NO:5:	
	(i)	SEQUE	NCE CHARACTERIST	CICS:
		(A)	LENGTH:	15
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUE	NCE DESCRIPTION:	SEQ ID NO:5:
		TTTTT	FTTTT TTTTT 15	
(2)	INFOR	MATION F	OR SEQ ID NO:6:	
	(i)	SEQUE	NCE CHARACTERIST	ICS:
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		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)		ICE DESCRIPTION:	
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(2)			OR SEQ ID NO:7:	
	(i)		CE CHARACTERIST	ICS:
		(A)	LENGTH:	24
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUEN	CE DESCRIPTION:	SEQ ID NO:7:
		GTAGTG	ATGC CATGTAACTG	AATC 24
(2)			OR SEQ ID NO:8:	
	(i)	SEQUEN	CE CHARACTERISTI	CS:
		(A)	LENGTH:	9
		(B)	TYPE:	amino acid
		(C)		single
		(D)	TOPOLOGY:	linear
	(xi)		CE DESCRIPTION:	
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5

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                (B)
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                (C)
                       STRANDEDNESS: double
                (D)
                       TOPOLOGY:
                                      linear
                SEQUENCE DESCRIPTION: SEQ ID NO:9:
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 CGCTGGGTCC CCTCTCCCCT GGCGCCCTGC CCCGACCTGC GCAAGCACAG GACGTCGTGG 180
 ACCTGGACTT CTTCACCCAG GAGCCGCTGC ACCTGGTGAG CCCCTCGTTC CTGTCCGTCA 240
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 CTCAAGTCAA CCAGGATATT TGCAAATATG GATCCATCCC TCCTGATGTG GAGGAGAAGT 480
 TACGGTTGGA ATGGCCCTAC CAGGAGCAAT TGCTACTCCG AGAACACTAC CAGAAAAAGT 540
 TCAAGAACAG CACCTACTCA AGAAGCTCTG TAGATGTGCT ATACACTTTT GCAAACTGCT 600
 CAGGACTGGA CTTGATCTTT GGCCTAAATG CGTTATTAAG AACAGCAGAT TTGCAGTGGA 660
ACAGTTCTAA TGCTCAGTTG CTCCTGGACT ACTGCTCTTC CAAGGGGTAT AACATTTCTT 720
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TTTCATCTGT GCAAAAAGTT TTCCAGGTGG TTGAGAGCAC CAGGCCTGGC AAGAAGGTCT 1080
GGTTAGGAGA AACAAGCTCT GCATATGGAG GCGGAGCGCC CTTGCTATCC GACACCTTTG 1140
CAGCTGGCTT TATGTGGCTG GATAAATTGG GCCTGTCAGC CCGAATGGGA ATAGAAGTGG 1200
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CTTTACCTGA TTATTGGCTA TCTCTTCTGT TCAAGAAATT GGTGGGCACC AAGGTGTTAA 1320
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CTGACAATCC AAGGTATAAA GAAGGAGATT TAACTCTGTA TGCCATAAAC CTCCATAACG 1440
TCACCAAGTA CTTGCGGTTA CCCTATCCTT TTTCTAACAA GCAAGTGGAT AAATACCTTC 1500
TAAGACCTTT GGGACCTCAT GGATTACTTT CCAAATCTGT CCAACTCAAT GGTCTAACTC 1560
TAAAGATGGT GGATGATCAA ACCTTGCCAC CTTTAATGGA AAAACCTCTC CGGCCAGGAA 1620
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(2)
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                     LENGTH:
                                543
               (B)
                      TYPE:
                                     amino acid
               (C)
                     STRANDEDNESS: single
                      TOPOLOGY:
               (D)
                                    linear
       (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
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                                    10
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Ala Gln Ala Gln Asp Val Val Asp Leu Asp Phe Phe Thr Gln Glu Pro
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Leu His Leu Val Ser Pro Ser Phe Leu Ser Val Thr Ile Asp Ala Asn

55

WO 00/52178

Le 6		a Ti	hr A	ap P		rg Pl 70	he Le	u Il	e Le	u Le 7		y Se:	r Pro	o Ly	s Leu 80
Ar	g Th	r L	eu A		rg G] 85	ly Le	eu Se	r Pr	o Al 9		r Leu	ı Arç	g Phe	e G1; 9:	y Gly
Th	r Ly	s Tì		sp Pl	ne Le	eu Il	e Ph	e As 10	-	o Ly	s Lys	Glu	Ser 110		. Phe
Gla	ı Gl	u Ar 11		er Ty	r Tr	p Gl	n Se 12		n Va	l Ası	ı Gln	As _F		Cyr	. Lys
Туз	13		r I	le Pi	o Pr	o As 13		l Gl	u Gli	Lys	140		Leu	Glu	Trp
Pro		r Gl	n Gl	.u G1	n Le 15		u Lei	ı Arç	g Glu	1 His	Tyr	Gln	Lys	Lys	Phe 160
Lys	Ası	a Se	r Th	r Ty 16		r Ar	g Sei	Sei	r Val		Val	Leu	Туr	Thr	Phe
Ala	Ası	1 Су	8 Se		y Le	u Asj	ρ Le u	11e		Gly	Leu	Asn	Ala 190	Leu	Leu
Arg	Thr	19		p Le	u Gli	ı Tr	Asn 200		Ser	Asn	Ala	Gln 205	Leu	Leu	Leu
Asp	Tyr 210		s Se	r Se	r Lys	Gly 215	-	naA	Ile	Ser	Trp 220	Glu	Leu	Gly	Asn
Glu 225	Pro	Ası	ı Se	r Pho	230		Lys	Ala	Авр	Ile 235	Phe	Ile	Asn	Gly	Ser 240
Gln	Leu	Gl	r Gl) Asp 245		Ile	Gln	Leu	His 250	Lys	Leu	Leu	Arg	<i>L</i> ys 255	Ser
Thr	Phe	Lys	260		Lys	Leu	Tyr	Gly 265	Pro	Asp	Val	Gly	Gln 270	Pro	Arg
Arg	ГЛа	Thr 275		Lys	Met	Leu	Lys 280	Ser	Phe	Leu	Lys	Ala 285	Gly	Gly	Glu
Val	Ile 290	Asp	Ser	Val	Thr	Trp 295	His	His	Tyr	Tyr	Leu 300	naA	Gly	Arg	Thr
Ala 305	Thr	Arg	Glu	Asp	Phe 310	Leu	Asn	Pro	Asp	Val 315	Leu .	Asp	Ile	Phe	11e 320
Ser	Ser	Val	Gln	Lys 325	Val	Phe	Gln	Val	Val 330	Glu	Ser'	Thr .	_	Pro 335	Gly
Lys	Lys	Val	Trp		Gly	Glu		Ser		Ala	Tyr (Gly (Gly	Ala

Pro Leu Ser Asp Thr Phe Ala Ala Gly Phe Met Trp Leu Asp Lys 355 360 365

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Le	u Gl 37		eu s	Ser	Ala	Arg	у Ме 37		y 11	e Gl	u Va	1 Va 38		t Arg	g Gl:	n Val	
Ph 38		e G	Ly #	Ala	Gly	Ası 390	-	r Hi	s Le	u Va	1 As 39	-	u Ası	n Phe	e Ası	Pro 400	
Le	u Pr	o As	p T	yr	Trp 405	Leu	Se:	r Le	ı Le	u Ph 41	-	s Ly:	s Let	val	G1 ₃	/ Thr	
Ly	s Va	l Le	eu M	let 420		Ser	Va.	l Glr		y Se. 25	r Ly	B Arg	a Arg	Lys 43		a Arg	
Va:	1 ту	r Le		is	Сув	Thr	Ası	440		p Ası	n Pro	o Arg	7 Tyr 445	-	Glu	Gly	
laA	450		r L	eu	Tyr	Ala	11e 455		Lei	a Hi	s Asr	1 Val		Lys	Туг	Leu	
Arg 469		ı Pr	о Т	yr	Pro	Phe 470	Ser	Asn	Lys	Glr	475	_	Lys	Tyr	Leu	Leu 480	
Arg	Pro	Le	u G		Pro 485	His	Gly	Leu	Let	Ser 490		Ser	Val	Gln	Leu 495		
Gly	Leu	Th		eu 00	Lys	Met	Val	Asp	Asp 505		Thr	Leu	Pro	Pro 510	Leu	Met	
Glu	Lys	Pro		eu .	Arg	Pro	Gly	Ser 520	Ser	Leu	Gly	Leu	Pro 525	Ala	Phe	Ser	
Tyr	Ser 530		e Pł	ıe '	Val	Ile	Arg 535	Asn	Ala	Lys	Val	Ala 540	Ala	Сув	Ile 543		
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					B) C)		YPE: TRAN	DEDN	ESS:			.c ac	10				
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		(xi)	S	EQUE	NCE	DESC	CRIPT	NOI	: SE	Q ID	10:	11:				
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TCT	CCG	CTG	CG	CG	GC 3	AGC	TGG	CGG	GGG	GAG	CAG	CCA	GGT	GAG	CCC	AAG	62
ATG	CTG	CTG	CG	c 1	CG A	AG	сст	GCG	CTG	CCG	CCG	CCG	CTG	ATG	CTG	CTG	110
Met	Leu	Leu	Ar	g S	er I	ys	Pro	Ala	Leu	Pro	Pro	Pro	Leu	Met	Leu	Leu	
					5					10					15		
CTC	CTG	GGG	CC	G C	TG G	GT (CCC	CTC	TCC	CCT	GGC	GCC	CTG	ccc	CGA	CCT	158
Leu	Leu	Gly	Pro		eu G	ly 1	Pro	Leu	Ser 25	Pro	Gly	Ala	Leu	Pro 30	Arg	Pro	
GCG	CAA	GCA	CA	G G	AC G	TC (STG	GAC	CTG	GAC	TTC	TTC	ACC	CAG	GAG	CCG	206
Ala	Gln		Gl	n A	sp V	al V	/al		Leu	Asp	Phe	Phe		Gln	Glu	Pro	
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CT	G CA	C CI	rg g	TG A	GC C	CC TC	G TI	c ci	G TC	C G1	C AC	C AT	T GA	C GC	C AAC	254
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	50					5	5				6	0				
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		a Th	r A	sp Pi		_	e Le	u Il	e Le			y Se	r Pr	o Ly	s Leu	
6	5				7	70				7	5				80	
															r GGC	
Arg	g Th	r Le	u Al			y Le	ı Se	r Pr		_	r Le	ı Arg	g Phe	_	Gly	
				85	•				9	0				9:	5	
200	. nn	7 70	2 62		·	·										
															TTT Phe	398
1111	. шу.	5 141	10		e be	u 110	PILE	10!		о цу	в пу	3 GIL			Pne	
			10					10:	,				110	,		
GAA	GAC	DA F	א א	т та	с та	G CA	י יייריין	ר כאז	CT(~ AA	~ CAC	יאם ב	י איי	TOO	ממה י	446
						p Glr										440
		11		,		p 01.	120					125		. Cya	, шуа	
TAT	GGZ	TC	CAT	c cc	T CC	T GAT	GTG	GAG	GAG	AA C	TTA	CGG	TTG	GAA	TGG	494
						o Asp										
	130)				135					140				_	
ccc	TAC	CAC	GA	G CA	A TT	G CTA	CTC	CGA	GAA	CAC	TAC	CAG	AAA	AAG	TTC	542
Pro	Tyr	Glr	ı Gl	u Gli	a Let	ı Leu	Leu	Arg	Glu	His	Туг	Gln	Lys	Lys	Phe	
145					150)				155					160	
						A AGA										590
Lys	Asn	Ser	Th:			Arg	Ser	Ser		_	Val	Leu	Tyr	Thr	Phe	
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						GAC										638
MIG	ABII	Cys	180		rec	qeA ı	Leu		Pne	GIY	Leu	Asn		Leu	Leu	
			100	,				185					190			
AGA	ACA	GCA	GAT	י ייייי	CAC	TGG	ממ	ACT	тст	λλΤ	CCT	CAG	TTTC	CTC	CTC	686
						Trp										000
		195	_				200	-				205		Dou	LCu	
GAC	TAC	TGC	TCI	TCC	AAG	GGG	TAT	AAC	ATT	TCT	TGG	GAA	CTA	GGC	AAT	734
						Gly										
	210					215					220					
GAA	CCT	AAC	AGT	TTC	CTT	AAG	AAG	GCT	GAT	ATT	TTC	ATC	AAT	GGG	TCG	782
Glu	Pro	Asn	Ser	Phe	Leu	Lys	Lys	Ala	Asp	Ile	Phe	Ile	Asn	Gly	Ser	
225					230					235					240	
						ATT										830
Gln	Leu	Gly	Glu		Tyr	Ile	Gln	Leu		ГÀа	Leu	Leu	Arg	-	Ser	
				245					250					255		
																
															CGA	878
inr	۲ne	ьys	Asn	Ala	гав	Leu	Tyr	Gly	Pro	qeA	Val	Gly	Gln	Pro	Arg	

7

AGA AAG ACG GCT AAG ATG CTG AAG AGC TTC CTG AAG GCT GGT GGA GAA 926
Arg Lys Thr Ala Lys Met Leu Lys Ser Phe Leu Lys Ala Gly Gly Glu
275 280 285

- GTG ATT GAT TCA GTT ACA TGG CAT CAC TAC TAT TTG AAT GGA CGG ACT 974

 Val Ile Asp Ser Val Thr Trp His His Tyr Tyr Leu Asn Gly Arg Thr

 290 295 300
- GCT ACC AGG GAA GAT TTT CTA AAC CCT GAT GTA TTG GAC ATT TTT ATT 1022
 Ala Thr Arg Glu Asp Phe Leu Asn Pro Asp Val Leu Asp Ile Phe Ile
 305 310 315 320
- TCA TCT GTG CAA AAA GTT TTC CAG GTG GTT GAG AGC ACC AGG CCT GGC 1070 Ser Ser Val Gln Lys Val Phe Gln Val Val Glu Ser Thr Arg Pro Gly 325 330 335
- AAG AAG GTC TGG TTA GGA GAA ACA AGC TCT GCA TAT GGA GGC GGA GCG 1118 Lys Lys Val Trp Leu Gly Glu Thr Ser Ser Ala Tyr Gly Gly Gly Ala 340 345 350
- CCC TTG CTA TCC GAC ACC TTT GCA GCT GGC TTT ATG TGG CTG GAT AAA 1166
 Pro Leu Leu Ser Asp Thr Phe Ala Ala Gly Phe Met Trp Leu Asp Lys
 355 360 365
- TTG GGC CTG TCA GCC CGA ATG GGA ATA GAA GTG GTG ATG AGG CAA GTA 1214
 Leu Gly Leu Ser Ala Arg Met Gly Ile Glu Val Val Met Arg Gln Val
 370 375 380
- TTC TTT GGA GGA GGA AAC TAC CAT TTA GTG GAT GAA AAC TTC GAT CCT 1262
 Phe Phe Gly Ala Gly Asn Tyr His Leu Val Asp Glu Asn Phe Asp Pro
 385 390 395 400
- TTA CCT GAT TAT TGG CTA TCT CTT CTG TTC AAG AAA TTG GTG GGC ACC 1310 Leu Pro Asp Tyr Trp Leu Ser Leu Leu Phe Lys Lys Leu Val Gly Thr 405 410 415
- AAG GTG TTA ATG GCA AGC GTG CAA GGT TCA AAG AGA AGG AAG CTT CGA 1358 Lys Val Leu Met Ala Ser Val Gln Gly Ser Lys Arg Arg Lys Leu Arg 420 425 430
- GTA TAC CTT CAT TGC ACA AAC ACT GAC AAT CCA AGG TAT AAA GAA GGA 1406
 Val Tyr Leu His Cys Thr Asn Thr Asp Asn Pro Arg Tyr Lys Glu Gly
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 440
 445
- GAT TTA ACT CTG TAT GCC ATA AAC CTC CAT AAC GTC ACC AAG TAC TTG 1454
 Asp Leu Thr Leu Tyr Ala Ile Asn Leu His Asn Val Thr Lys Tyr Leu
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 450
 460
- CGG TTA CCC TAT CCT TTT TCT AAC AAG CAA GTG GAT AAA TAC CTT CTA 1502
 Arg Leu Pro Tyr Pro Phe Ser Asn Lys Gln Val Asp Lys Tyr Leu Leu
 465 470 475 480
- AGA CCT TTG GGA CCT CAT GGA TTA CTT TCC AAA TCT GTC CAA CTC AAT 1550 Arg Pro Leu Gly Pro His Gly Leu Leu Ser Lys Ser Val Gln Leu Asn 485 490 495

GGT CTA ACT CTA AAG ATG GTG GAT GAT CAA ACC TTG CCA CCT TTA ATG 1598

8

Gly Leu Thr Leu Lys Met Val Asp Asp Gln Thr Leu Pro Pro Leu Met 505

GAA AAA CCT CTC CGG CCA GGA AGT TCA CTG GGC TTG CCA GCT TTC TCA 1646 Glu Lys Pro Leu Arg Pro Gly Ser Ser Leu Gly Leu Pro Ala Phe Ser 520

TAT AGT TTT TTT GTG ATA AGA AAT GCC AAA GTT GCT GCT TGC ATC TGA 1694 Tyr Ser Phe Phe Val Ile Arg Asn Ala Lys Val Ala Ala Cys Ile 535 540

AAA TAA AAT ATA CTA GTC CTG ACA CTG 1721

- INFORMATION FOR SEQ ID NO:12: (2)
 - (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 824

(B) TYPE: nucleic acid

- (C) STRANDEDNESS: double
- (D) TOPOLOGY:
 - linear
- SEQUENCE DESCRIPTION: SEQ ID NO:12 (xi)

CTGGCAAGAA GGTCTGGTTG GGAGAGACGA GCTCAGCTTA CGGTGGCGGT GCACCCTTGC 60 TGTCCAACAC CTTTGCAGCT GGCTTTATGT GGCTGGATAA ATTGGGCCTG TCAGCCCAGA 120 TGGGCATAGA AGTCGTGATG AGGCAGGTGT TCTTCGGAGC AGGCAACTAC CACTTAGTGG 180 ATGAAAACTT TGAGCCTTTA CCTGATTACT GGCTCTCTCT TCTGTTCAAG AAACTGGTAG 240 GTCCCAGGGT GTTACTGTCA AGAGTGAAAG GCCCAGACAG GAGCAAACTC CGAGTGTATC 300 TCCACTGCAC TAACGTCTAT CACCCACGAT ATCAGGAAGG AGATCTAACT CTGTATGTCC 360 TGAACCTCCA TAATGTCACC AAGCACTTGA AGGTACCGCC TCCGTTGTTC AGGAAACCAG 420 TGGATACGTA CCTTCTGAAG CCTTCGGGGC CGGATGGATT ACTTTCCAAA TCTGTCCAAC 480 TGAACGGTCA AATTCTGAAG ATGGTGGATG AGCAGACCCT GCCAGCTTTG ACAGAAAAAC 540 CTCTCCCCGC AGGAAGTGCA CTAAGCCTGC CTGCCTTTTC CTATGGTTTT TTTGTCATAA 600 GAAATGCCAA AATCGCTGCT TGTATATGAA AATAAAAGGC ATACGGTACC CCTGAGACAA 660 AAGCCGAGGG GGGTGTTATT CATAAAACAA AACCCTAGTT TAGGAGGCCA CCTCCTTGCC 720 GAGTTCCAGA GCTTCGGGAG GGTGGGGTAC ACTTCAGTAT TACATTCAGT GTGGTGTTCT 780 CTCTAAGAAG AATACTGCAG GTGGTGACAG TTAATAGCAC TGTG 824

- (2) INFORMATION FOR SEQ ID NO:13:
 - SEQUENCE CHARACTERISTICS: (i)

LENGTH:

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY:

- SEQUENCE DESCRIPTION: SEQ ID NO:13 (xi)
- GGGAAAGCGA GCAAGGAAGT AGGAGAGAGC CGGGCAGGCG GGGCGGGGTT GGATTGGGAG 60 CAGTGGGAGG GATGCAGAAG AGGAGTGGGA GGGATGGAGG GCGCAGTGGG AGGGGTGAGG 120 AGGCGTAACG GGGCGGAGGA AAGGAGAAAA GGGCGCTGGG GCTCGGCGGG AGGAAGTGCT 180 AGAGCTCTCG ACTCTCCGCT GCGCGGCAGC TGGCCGGGGG AGCAGCCAGG TGAGCCCAAG ATGCTGCTGC GCTCGAAGCC TGCGCTGCCG CCGCCGCTGA TGCTGCTGCT CCTGGGGCCG 300 CTGGGTCCCC TCTCCCCTGG CGCCCTGCCC CGACCTGCGC AAGCACAGGA CGTCGTGGAC 360 CTGGACTTCT TCACCCAGGA GCCGCTGCAC CTGGTGAGCC CCTCGTTCCT GTCCGTCACC 420 ATTGACGCCA ACCTGGCCAC GGACCCGCGG TTCCTCATCC TCCTGGGTTC TCCAAAGCTT 480 CGTACCTTGG CCAGAGGCTT GTCTCCTGCG TACCTGAGGT TTGGTGGCAC CAAGACAGAC 540 TTCCTAATTT TCGATCCCAA GAAGGAATCA ACCTTTGAAG AGAGAAGTTA CTGGCAATCT CAAGTCAACC AGGATATTTG CAAATATGGA TCCATCCCTC CTGATGTGGA GGAGAAGTTA CGGTTGGAAT GGCCCTACCA GGAGCAATTG CTACTCCGAG AACACTACCA GAAAAGGTTC 720 AAGAACAGCA CCTACTCAAG AAGCTCTGTA GATGTGCTAT ACACTTTTGC AAACTGCTCA 780

				-		
GGACTGGACT	TGATCTTTGG	CCTAAATGCG	TTATTAAGAA	CAGCAGATTT	GCAGTGGAAC	840
AGTTCTAATO	CTCAGTTGCT	CCTGGACTAC	TGCTCTTCCA	AGGGGTATAA	CATTTCTTGG	900
GAACTAGGCA	ATGAACCTAA	CAGTTTCCTT	AAGAAGGCTG	ATATTTTCAT	CAATGGGTCG	960
CAGTTAGGAG	AAGATTATAT	TCAATTGCAT	AAACTTCTAA	GAAAGTCCAC	CTTCAAAAAT	1020
GCAAAACTCT	ATGGTCCTGA	TGTTGGTCAG	CCTCGAAGAA	AGACGGCTAA	GATGCTGAAG	1080
AGCTTCCTGA	AGGCTGGTGG	AGAAGTGATT	GATTCAGTTA	CATGGCATCA	CTACTATTTG	1140
AATGGACGGA	CTGCTACCAG	GGAAGATTTT	CTAAACCCTG	ATGTATTGGA	CATTTTTATT	1200
TCATCTGTGC	AAAAAGTTTT	CCAGGTGGTT	GAGAGCACCA	GGCCTGGCAA	GAAGGTCTGG	1260
TTAGGAGAAA	CAAGCTCTGC	ATATGGAGGC	GGAGCGCCCT	TGCTATCCGA	CACCTTTGCA	1320
GCTGGCTTTA	TGTGGCTGGA	TAAATTGGGC	CTGTCAGCCC	GAATGGGAAT	AGAAGTGGTG	1380
ATGAGGCAAG	TATTCTTTGG	AGCAGGAAAC	TACCATTTAG	TGGATGAAAA	CTTCGATCCT	1440
TTACCTGATT	ATTGGCTATC	TCTTCTGTTC	AAGAAATTGG	TGGGCACCAA	GGTGTTAATG	1500
GCAAGCGTGC	AAGGTTCAAA	GAGAAGGAAG	CTTCGAGTAT	ACCTTCATTG	CACAAACACT	1560
GACAATCCAA	GGTATAAAGA	AGGAGATTTA	ACTCTGTATG	CCATAAACCT	CCATAACGTC	1620
ACCAAGTACT	TGCGGTTACC	CTATCCTTTT	TCTAACAAGC	AAGTGGATAA	ATACCTTCTA	1680
AGACCTTTGG	GACCTCATGG	ATTACTTTCC	AAATCTGTCC	AACTCAATGG	TCTAACTCTA	1740
AAGATGGTGG	ATGATCAAAC	CTTGCCACCT	TTAATGGAAA	AACCTCTCCG	GCCAGGAAGT	1800
	TGCCAGCTTT			TAAGAAATGC	CAAAGTTGCT	1860
GCTTGCATCT	GAAAATAAAA	TATACTAGTC	CTGACACTG			1899

(2) INFORMATION FOR SEQ ID NO:14:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 592

(B) TYPE: amino acid

(C) STRANDEDNESS: singl

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14

Met Glu Gly Ala Val Gly Gly Val Arg Arg Arg Asn Gly Ala Glu 5 10 Glu Arg Arg Lys Gly Arg Trp Gly Ser Ala Gly Gly Ser Ala Arg 20 25 Ala Leu Asp Ser Pro Leu Arg Gly Ser Trp Arg Gly Glu Gln Pro 35 40 Gly Glu Pro Lys Met Leu Leu Arg Ser Lys Pro Ala Leu Pro Pro 50 55 60 Pro Leu Met Leu Leu Leu Gly Pro Leu Gly Pro Leu Ser Pro 65 70 Gly Ala Leu Pro Arg Pro Ala Gln Ala Gln Asp Val Val Asp Leu 80 85 Asp Phe Phe Thr Gln Glu Pro Leu His Leu Val Ser Pro Ser Phe 95 100 105 Leu Ser Val Thr Ile Asp Ala Asn Leu Ala Thr Asp Pro Arg Phe 110 115 Leu Ile Leu Leu Gly Ser Pro Lys Leu Arg Thr Leu Ala Arg Gly 125 130 Leu Ser Pro Ala Tyr Leu Arg Phe Gly Gly Thr Lys Thr Asp Phe 140 145 Leu Ile Phe Asp Pro Lys Lys Glu Ser Thr Phe Glu Glu Arg Ser 160 155 Tyr Trp Gln Ser Gln Val Asn Gln Asp Ile Cys Lys Tyr Gly Ser 170 175 Ile Pro Pro Asp Val Glu Glu Lys Leu Arg Leu Glu Trp Pro Tyr 185 190 195 Gln Glu Gln Leu Leu Arg Glu His Tyr Gln Lys Lys Phe Lys 200 205 210 Asn Ser Thr Tyr Ser Arg Ser Ser Val Asp Val Leu Tyr Thr Phe

Ala Asn Cys Ser Gly Leu Asp Leu Ile Phe Gly Leu Asn Ala Leu Leu Arg Thr Ala Asp Leu Gln Trp Asn Ser Ser Asn Ala Gln Leu Leu Leu Asp Tyr Cys Ser Ser Lys Gly Tyr Asn Ile Ser Trp Glu Leu Gly Asn Glu Pro Asn Ser Phe Leu Lys Lys Ala Asp Ile Phe Ile Asn Gly Ser Gln Leu Gly Glu Asp Tyr Ile Gln Leu His Lys Leu Leu Arg Lys Ser Thr Phe Lys Asn Ala Lys Leu Tyr Gly Pro Asp Val Gly Gln Pro Arg Arg Lys Thr Ala Lys Met Leu Lys Ser Phe Leu Lys Ala Gly Gly Glu Val Ile Asp Ser Val Thr Trp His His Tyr Tyr Leu Asn Gly Arg Thr Ala Thr Arg Glu Asp Phe Leu Asn Pro Asp Val Leu Asp Ile Phe Ile Ser Ser Val Gln Lys Val Phe Gln Val Val Glu Ser Thr Arg Pro Gly Lys Lys Val Trp Leu 385. Gly Glu Thr Ser Ser Ala Tyr Gly Gly Gly Ala Pro Leu Leu Ser Asp Thr Phe Ala Ala Gly Phe Met Trp Leu Asp Lys Leu Gly Leu Ser Ala Arg Met Gly Ile Glu Val Val Met Arg Gln Val Phe Phe Gly Ala Gly Asn Tyr His Leu Val Asp Glu Asn Phe Asp Pro Leu Pro Asp Tyr Trp Leu Ser Leu Leu Phe Lys Lys Leu Val Gly Thr Lys Val Leu Met Ala Ser Val Gln Gly Ser Lys Arg Arg Lys Leu Arg Val Tyr Leu His Cys Thr Asn Thr Asp Asn Pro Arg Tyr Lys Glu Gly Asp Leu Thr Leu Tyr Ala Ile Asn Leu His Asn Val Thr Lys Tyr Leu Arg Leu Pro Tyr Pro Phe Ser Asn Lys Gln Val Asp Lys Tyr Leu Leu Arg Pro Leu Gly Pro His Gly Leu Leu Ser Lys Ser Val Gln Leu Asn Gly Leu Thr Leu Lys Met Val Asp Asp Gln Thr Leu Pro Pro Leu Met Glu Lys Pro Leu Arg Pro Gly Ser Ser Leu Gly Leu Pro Ala Phe Ser Tyr Ser Phe Phe Val Ile Arg Asn Ala Lys Val Ala Ala Cys Ile

(2) INFORMATION FOR SEQ ID NO:15:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1899
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double

											1 1				
				(D)			LOGY			inea					
		(xi	,	SEQU	JENCE	: DE:	SCRI	PTION	i: S	EQ I	D NO	:15			
														GGG	3
222	GCC	ם אכו	י ממי	CAR	СТВ	CC	CA(ם אפר	ccc	י פרא	GGC	ccc	e co	GGG	48
								A TGC							93
						-		G AGG					-		138
								l Arg							130
		. 017		. 142	_	٠.,	, , ,		10	9	7.01		,,,,	15	
				_											
GAA	AGG	AGA	AAA	GGG	CGC	TGC	GGG	TCG	GCG	GGA	GGA	AGT	GCT	AGA	183
								Ser							
	_		-	20	_	-			25		-			30	
GCT	CTC	GAC	TCT	CCG	CTG	CGC	GGC	AGC	TGG	CGG	GGG	GAG	CAG	CCA	228
Ala	Leu	Авр	Ser	Pro	Leu	Arg	Gly	Ser	Trp	Arg	Gly	Glu	Gln	Pro	
				35					40					45	
GGT	GAG	ccc	AAG	ATG	CTG	CTG	CGC	TCG	AAG	CCT	GCG	CTG	CCG	CCG	273
Gly	Glu	Pro	Lys	Met	Leu	Leu	Arg	Ser	Lув	Pro	Ala	Leu	Pro	Pro	
				50					55					60	
								CCG							318
Pro	Leu	Met	Leu		Leu	Leu	Gly	Pro		Gly	Pro	Leu	Ser		
				65					70					75	
000	ccc	ama	000	CC 3	com	000	CNN	CCN	C2.C	a.c	am-	ana	C2.C	CTC.	363
								GCA Ala							363
GIY	AIA	Leu	PIO	80	PIO	MIG	GIII	мта	85	ивр	Val	vai	жыр	90	
				80					65					30	
GAC	TTC	TTC	ACC	CAG	GAG	CCG	CTG	CAC	CTG	GTG	AGC	ccc	TCG	TTC	408
								His							
				95					100					105	
CTG	TCC	GTC	ACC	ATT	GAC	GCC	AAC	CTG	GCC	ACG	GAC	CÇG	CGG	TTC	453
Leu	Ser	Val	Thr	Ile	qaA	Ala	Asn	Leu	Ala	Thr	Asp	Pro	Arg	Phe	
				110					115					120	
CTC	ATC	CTC	CTG	GGT	TCT	CCA	AAG	CTT	CGT	ACC	TTG	GCC	AGA	GGC	498
Leu	Ile	Leu	Leu	Gly	Ser	Pro	Lys	Leu	Arg	Thr	Leu	Ala	Arg	Gly	
				125					130					135	
								GGT							543
Leu	Ser	Pro	Ala	-	Leu	Arg	Phe	Gly	-	Thr	ГÀе	Thr	Asp		
				140					145					150	
COLD	a mm	mmo	C N TT	ccc	220	220	C N N	TCA	200	man	~~~	CAC	202	NOT	E 0 0
								Ser							588
пец	116	FIIC	_	155	шув	Lys	GIU		160	FIIC	GLU	GIU	ura	165	
														-03	
TAC	TGG	CAA	TCT	CAA	GTC	AAC	CAG	GAT	АТТ	TGC	AAA	TAT	GGA	TCC	633
								Asp							
				170				-	175	·				180	
ATC	ССТ	CCT	GAT	GTG	GAG	GAG	AAG	TTA	CGG	TTG	GAA	TGG	ccc	TAC	678
Ile	Pro	Pro	Asp	Val	Glu	Glu	Lys	Leu .	Arg	Leu	Glu	Trp	Pro	Tyr	
				185					190					195	

CA	G	AG	CA	A TI	G CI	A CT	C CG	A GA	A CA	C TA	C CA	G AA	A AA	G TT	C AAG	723
Gl	n C	lu	Gl	n Le	u Le	u Le	u Ar	g Gl	u Hi	в Ту	r Gl	n Ly	s Ly	s Ph	e Lys	
					20	0				20	5				210	
															TTT	768
As	n S	er	Th	г Ту			g Se	r Se	r Val	L As	p Va	l Le	и ту	r Thi	r Phe	
					21	5				220	0				225	
															TTA	813
AI	a A	sn	Сує	s se			ı Ası	p Lei	1 Ile			y Lei	ı Ası	1 Ala	Leu	
					23	U				235	•				240	
тт	ב ב	GA	ACZ	GC	A GD.	י יייי	ב ראר	a mee	ב אאר	י אכי	, m.c.	י החי			TTG	858
								Trp								030
		- 5			24		. 0		, non	250		. ADI		GII	255	
CT	C (rg	GAC	TAC	TG	TCT	TCC	: AAG	GGG	TAT	' AAC	ATT	TCT	TGG	GAA	903
								Lys								
					260			_	_	265				•	270	
CTA	G	GC	AAT	GAJ	CCI	AAC	AGT	TTC	CTT	AAG	AAG	GCT	' GAI	ATT	TTC	948
Lev	ı G	lу	Asn	Glu	Pro	Asn	Ser	Phe	Leu	Lys	Lys	Ala	Asp	Ile	Phe	
					275	,				280					285	
		_														
								GAA								993
Ile	: As	m	Gly	Ser			Gly	Glu	Asp	-	Ile	Gln	Leu	His	-	
					290					295					300	
CTT	י ריז	מי	aca	A D C	TOO	N.C.C	TTC	AAA	3 3 T	CCN	222		m> m		aam	
								Lys								1038
			3	-,-	305		1110	273		310	Dy 0	Deu	171	GIY	315	
GAT	GI	T (GGT	CAG	CCT	CGA	AGA	AAG	ACG	GCT	AAG	ATG	CTG	AAG	AGC	1083
Asp	٧a	1 (Gly	Gln	Pro	Arg	Arg	Lys	Thr	Ala	Lys	Met	Leu	Lys	Ser	
					320					325				-	330	
TTC	CT	G A	AAG	GCT	GGT	GGA	GAA	GTG	ATT	GAT	TCA	GTT	ACA	TGG	CAT	1128
Phe	Le	u 1	Lys	Ala	Gly	Gly	Glu	Val	Ile	qaA	Ser	Val	Thr	Trp	His	
					335					340					345	
								ACT								1173
UIB	19	E 1	уr	Leu		GIA	Arg	Thr			Arg	Glu	Asp	Phe		
					350					355					360	
AAC	CC	го	AT	GTA	ттс	GAC	тта	TTT	ልጥጥ	ፈጋጥ	тст	GTG	CAA	222	CTT	1218
								Phe								1216
			•		365					370				-,.	375	
TTC	CAG	3 G	TG	GTT	GAG	AGC	ACC	AGG	CCT	GGC	AAG	AAG	GTC	TGG	TTA	1263
Phe	Gl	a V	al	Val	Glu	Ser	Thr	Arg	Pro	Gly	Lys	Lys	Val	Trp	Leu	
					380					385				_	390	
٠																
								GGA (1308
Gly	Gl	1 T	hr	Ser		Ala	Tyr	Gly (Ala	Pro	Leu	Leu	Ser	
					395				•	400					405	

13

GAC ACC TTT GCA GCT GGC TTT ATG TGG CTG GAT AAA TTG GGC CTG Asp Thr Phe Ala Ala Gly Phe Met Trp Leu Asp Lys Leu Gly Leu 410 415 TCA GCC CGA ATG GGA ATA GAA GTG GTG ATG AGG CAA GTA TTC TTT Ser Ala Arg Met Gly Ile Glu Val Val Met Arg Gln Val Phe Phe 425 430 GGA GCA GGA AAC TAC CAT TTA GTG GAT GAA AAC TTC GAT CCT TTA Gly Ala Gly Asn Tyr His Leu Val Asp Glu Asn Phe Asp Pro Leu 440 445 CCT GAT TAT TGG CTA TCT CTT CTG TTC AAG AAA TTG GTG GGC ACC Pro Asp Tyr Trp Leu Ser Leu Leu Phe Lys Lys Leu Val Gly Thr AAG GTG TTA ATG GCA AGC GTG CAA GGT TCA AAG AGA AGG AAG CTT Lys Val Leu Met Ala Ser Val Gln Gly Ser Lys Arg Arg Lys Leu 475 CGA GTA TAC CTT CAT TGC ACA AAC ACT GAC AAT CCA AGG TAT AAA Arg Val Tyr Leu His Cys Thr Asn Thr Asp Asn Pro Arg Tyr Lys 490 GAA GGA GAT TTA ACT CTG TAT GCC ATA AAC CTC CAT AAC GTC ACC Glu Gly Asp Leu Thr Leu Tyr Ala Ile Asn Leu His Asn Val Thr 500 505 AAG TAC TTG CGG TTA CCC TAT CCT TTT TCT AAC AAG CAA GTG GAT Lys Tyr Leu Arg Leu Pro Tyr Pro Phe Ser Asn Lys Gln Val Asp 515 520 AAA TAC CTT CTA AGA CCT TTG GGA CCT CAT GGA TTA CTT TCC AAA Lys Tyr Leu Leu Arg Pro Leu Gly Pro His Gly Leu Leu Ser Lys 530 535 TCT GTC CAA CTC AAT GGT CTA ACT CTA AAG ATG GTG GAT GAT CAA Ser Val Gln Leu Asn Gly Leu Thr Leu Lys Met Val Asp Asp Gln 545 550 ACC TTG CCA CCT TTA ATG GAA AAA CCT CTC CGG CCA GGA AGT TCA Thr Leu Pro Pro Leu Met Glu Lys Pro Leu Arg Pro Gly Ser Ser 565 CTG GGC TTG CCA GCT TTC TCA TAT AGT TTT TTT GTG ATA AGA AAT 1848 Leu Gly Leu Pro Ala Phe Ser Tyr Ser Phe Phe Val Ile Arg Asn 575 580 GCC AAA GTT GCT GCT TGC ATC TGA AAA TAA AAT ATA CTA GTC CTG 1893 Ala Lys Val Ala Ala Cys Ile 590 592 ACA CTG 1899 INFORMATION FOR SEO ID NO:16:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 594

TYPE:

nucleic acid

(B)

PCT/US00/03542 WO 00/52178

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14
                (C)
                       STRANDEDNESS: double
                (D)
                       TOPOLOGY:
                                      linear
                SEQUENCE DESCRIPTION: SEO ID NO:16
 ATTACTATAG GGCACGCGTG GTCGACGGCC CGGGCTGGTA TTGTCTTAAT GAGAAGTTGA 60
 TAAAGAATTT TGGGTGGTTG ATCTCTTTCC AGCTGCAGTT TAGCGTATGC TGAGGCCAGA 120
 TTTTTCAGG CAAAAGTAAA ATACCTGAGA AACTGCCTGG CCAGAGGACA ATCAGATTTT 180
 GGCTGGCTCA AGTGACAAGC AAGTGTTTAT AAGCTAGATG GGAGAGGAAG GGATGAATAC 240
 TCCATTGGAG GCTTTACTCG AGGGTCAGAG GGATACCCGG CGCCATCAGA ATGGGATCTG 300
 GGAGTCGGAA ACGCTGGGTT CCCACGAGAG CGCGCAGAAC ACGTGCGTCA GGAAGCCTGG 360
 TCCGGGATGC CCAGCGCTGC TCCCCGGGCG CTCCTCCCCG GGCGCTCCTC CCCAGGCCTC 420
 CCGGGCGCTT GGATCCCGGC CATCTCCGCA CCCTTCAAGT GGGTGTGGGT GATTTCGTAA 480
 GTGAACGTGA CCGCCACCGG GGGGAAAGCG AGCAAGGAAG TAGGAGAGAG CCGGGCAGGC 540
 GGGGCGGGT TGGATTGGGA GCAGTGGGAG GGATGCAGAA GAGGAGTGGG AGGG
 (2)
        INFORMATION FOR SEQ ID NO:17:
        (i)
               SEQUENCE CHARACTERISTICS:
                      LENGTH: 21
               (A)
               (B)
                       TYPE:
                                      nucleic acid
               (C)
                      STRANDEDNESS: single
               (D)
                      TOPOLOGY:
                                     linear
        (xi)
               SEQUENCE DESCRIPTION: SEQ ID NO:17
              CCCCAGGAGC AGCAGCATCA G 21
(2)
       INFORMATION FOR SEQ ID NO:18:
              SEQUENCE CHARACTERISTICS:
        (i)
               (A)
                      LENGTH:
                                 21
               (B)
                      TYPE:
                                     nucleic acid
               (C)
                      STRANDEDNESS: single
               (D)
                      TOPOLOGY:
                                     linear
     · (xi)
              SEQUENCE DESCRIPTION: SEQ ID NO:18
              AGGCTTCGAG CGCAGCAGCA T 21
(2)
       INFORMATION FOR SEQ ID NO:19:
              SEQUENCE CHARACTERISTICS:
               (A)
                      LENGTH:
                                     22
               (B)
                      TYPE:
                                     nucleic acid
              (C)
                     STRANDEDNESS: single
              (D)
                      TOPOLOGY:
                                   linear
              SEQUENCE DESCRIPTION: SEQ ID NO:19
       (xi)
             GTAATACGAC TCACTATAGG GC 22
(2)
       INFORMATION FOR SEO ID NO:20:
              SEQUENCE CHARACTERISTICS:
              (A)
                     LENGTH:
                                    19
              (B)
                     TYPE:
                                    nucleic acid
              (C)
                     STRANDEDNESS: single
              (D)
                     TOPOLOGY:
            SEQUENCE DESCRIPTION: SEQ ID NO:20
       (xi)
             ACTATAGGGC ACGCGTGGT 19
(2)
       INFORMATION FOR SEQ ID NO:21:
              SEQUENCE CHARACTERISTICS:
              (A)
                     LENGTH:
              (B)
                     TYPE:
                                    nucleic acid
              (C)
                     STRANDEDNESS: single
                     TOPOLOGY:
              (D)
                                    linear
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				15
	(xi)	SEQU	JENCE DESCRIPTION	: SEQ ID NO:2
		CTTGG	GCTCA CCTGGCTGCT	C 21
(2)	INFO	MATION	FOR SEQ ID NO: 2:	2:
	(i)		ENCE CHARACTERIS	
	,	(A)	LENGTH:	23
		(B)	TYPE:	nucleic acid
		(C)		
			STRANDEDNESS:	-
		(D)	TOPOLOGY:	linear
	(xi)		ENCE DESCRIPTION	
		AGCTC	TGTAG ATGTGCTATA	CAC 23
(2)			FOR SEQ ID NO:23	
	(i)	SEQU	ENCE CHARACTERIS	rics:
		(A)	LENGTH:	22
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUI	ENCE DESCRIPTION	SEO ID NO:23
			TTAGC CGTCTTTCTT	
(2)	INFOR	MATION	FOR SEQ ID NO:24	:
	(i)		ENCE CHARACTERIST	
		(A)	LENGTH:	23
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	
				_
		(D)	TOPOLOGY:	linear
	(xi)		NCE DESCRIPTION:	SEQ ID NO:24
GAGCA	GCCAG GT	GAGCCCA	A GAT 23	
(2)			FOR SEQ ID NO:25	
	(i)		NCE CHARACTERIST	
		(A)	LENGTH:	23
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUE	NCE DESCRIPTION:	SEQ ID NO:25
TTCGA	rccca aga	AAGGAAT	C AAC 23	
(2)	INFORM	ATION F	OR SEQ ID NO:26:	
	(i)	SEQUE	NCE CHARACTERIST	ICS:
		(A)	LENGTH:	23
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)		NCE DESCRIPTION:	
ACCTCT	GTAG ATO			SEQ ID NO:26
AGCICI	GIAG AIG	, IGCIAI	A CAC 23	
(2)	TNFORM	ATTON F	OR SEQ ID NO:27:	
127	(i)		CE CHARACTERISTI	ce.
	(1)	(A)		
			LENGTH:	24
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)		CE DESCRIPTION:	SEQ ID NO:27
TCAGAT	GCAA GCA	GCAACTT	TGGC 24	

				16
(2)	INFOR	MATION	FOR SEQ ID NO:28	3:
	(i)	SEQU	ENCE CHARACTERIS	TICS:
		(A)	LENGTH:	22
		(B)	TYPE:	nucleic acid
		·(C)	STRANDEDNESS:	
		(D)	TOPOLOGY:	linear
	(xi)		ENCE DESCRIPTION	
GCATC			TT CG 22	. 529 15 16.20
(2)	INFOR	MATION	FOR SEQ ID NO:29	·:
	(i)		NCE CHARACTERIS	
		(A)	LENGTH:	24
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUE	NCE DESCRIPTION:	
GTAGTO	GATGC CA		G AATC 24	-
(2)	INFORM	ATION I	FOR SEQ ID NO:30	:
	(i)	SEQUE	NCE CHARACTERIST	'ICS:
		(A)	LENGTH:	22
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	single
		(D)	TOPOLOGY:	linear
	(xi)	SEQUE	NCE DESCRIPTION:	SEQ ID NO:30
AGGCAC	CCTA GAG	GATGTTC	C AG 22	
(2)			OR SEQ ID NO:31:	
	(i)		NCE CHARACTERIST	
		(A)	LENGTH:	24
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	-
		(D)	TOPOLOGY:	linear
CAACAM	(xi)		ICE DESCRIPTION:	SEQ ID NO:31
GAAGAT.	ricr Gri	TCCATGA	CGTG 24	
(2)	TNEODM	ATTON E	OR SEQ ID NO:32:	
(2)	(i)			
	(1)	(A)	CE CHARACTERISTI LENGTH:	25
		(B)	TYPE:	nucleic acid
		(C)		single
		(D)	TOPOLOGY:	linear
	(xi)		CE DESCRIPTION:	
CCACACT	GAA TGT.			01Q 1D NO.32
(2)	INFORMA	TION FO	R SEQ ID NO:33:	
	(i)		CE CHARACTERISTI	CS:
		(A)	LENGTH:	22
		(B)	TYPE:	nucleic acid
		(C)	STRANDEDNESS:	
		(D)	TOPOLOGY:	linear
	(xi)	SEQUENC	CE DESCRIPTION:	
CGAAGCT	CTG GAAG			
(2)	INFORMA	TION FO	R SEQ ID NO:34:	
			E CHARACTERISTIC	CS:

(A)

LENGTH:

17 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear SEQUENCE DESCRIPTION: SEQ ID NO:34 GCCAGCTGCA AAGGTGTTGG AC 22 INFORMATION FOR SEQ ID NO:35: (2) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear SEQUENCE DESCRIPTION: SEQ ID NO:35 AACACCTGCC TCATCACGAC TTC 23 INFORMATION FOR SEQ ID NO:36: (2) SEQUENCE CHARACTERISTICS: (A) LENGTH: (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear SEQUENCE DESCRIPTION: SEQ ID NO:36 GCCAGGCTGG CGTCGATGGT GA 22 (2) INFORMATION FOR SEQ ID NO:37: SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear SEQUENCE DESCRIPTION: SEO ID NO:37 GTCGATGGTG ATGGACAGGA AC 22 (2) INFORMATION FOR SEQ ID NO:38: SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 22 TYPE: nucleic acid (C) STRANDEDNESS: single TOPOLOGY: · linear (D) SEQUENCE DESCRIPTION: SEQ ID NO:38 GTAATACGAC TCACTATAGG GC 22 (2) INFORMATION FOR SEQ ID NO:39: SEQUENCE CHARACTERISTICS: (i) (A) LENGTH: 19 (B) TYPE: nucleic acid STRANDEDNESS: single (C) (D) TOPOLOGY: linear (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39 ACTATAGGGC ACGCGTGGT 19 (2) INFORMATION FOR SEQ ID NO:40: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27

(B)

(C)

TYPE:

TOPOLOGY:

STRANDEDNESS: single

nucleic acid

linear

18

SEQUENCE DESCRIPTION: SEQ ID NO:40 (xi) CCATCCTAAT ACGACTCACT ATAGGGC 27

(2) INFORMATION FOR SEO ID NO:41:

(i) SEQUENCE CHARACTERISTICS:

> (A) LENGTH: 23

(B) TYPE:

nucleic acid

STRANDEDNESS: single (C)

> TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41

ACTCACTATA GGGCTCGAGC GGC 23

(xi)

(2) INFORMATION FOR SEO ID NO:42:

(D)

SEQUENCE CHARACTERISTICS:

(A) LENGTH:

(B) TYPE: nucleic acid

100

150

200

250

300

350

400

500

550

600

700

750

800

900

950

1000

1050

1100 1150

1200

1250

1300 1350

1400

1450

1500

1550

1600

1650

1700

1750

1800

1850

1900

1950

2000

2050

2100

2150

(C) STRANDEDNESS: double

(D) TOPOLOGY:

linear SEQUENCE DESCRIPTION: SEQ ID NO:42

GGATCTTGGC TCACTGCAAT CTCTGCCTCC CATGCAATTC TTATGCATCA GCCTCCTGAG TAGCTTGGAT TATAGGTCTG CGCCACCACT CCTGGCTACA CCATGTTGCC CAGGCTGGTC TTGAACTCTT GGGCTCTAGT GATCCACCCG CCTTGGCCTC CCAAAGTGCT GGGATTACAG GTGTGAGCCA TCACACCCGG CCCCCCGTTT CCATATTAGT AACTCACATG TAGACCACAA GGATGCACTA

TTTAGAAAAC TTGCAATGGT CCACTTTTCA AATCACCCAA ACATGTTAAA GAAATTGGTA TGACTGGGCA TGGCACAGTG GCTCATGCCT GCAATCCTAG CATTTTGTGA GGCTGAGACG GGCAGATCAC GAGGTCAGGA GATTGAGACC ATCCTGACAG ACATGGTGAA ATCCCATCTC TACTAAAAAT ACAAAACAAT TAGCCGGGGG TGATGGCAGG CCCCTGTAGT CCCAGCTACT CGGGAGGCTG AGGCAGGAGA ATGGCGTGAA TCCAGGAGGC AGAGCTTGCA GTGAGCCGAG

ATGGTGCCAC TGCACTCCAG CCTGGGCGAC AGAGCGAGAC TCCGTCTCAA AAAAAAAAA AAAGAAAGAA ATTGGTATGA CTGTTGACTC ACAACAGGAG TCAGGGGCAT GGGGTGGGGT GTAAGATTAA TGTCATGACA AATGTGGAAA AGAAACTTCT GTTTTTCCAA CTCCACGTCT GCTACCATAT TATTACACTC TTCTGGTAGT GTGGTGTTTA TGTGTGAATT TTTTTTCATA TGTATACAGT AATTGTAGGA TATGAACCTG ATTCTAGTTG CAAAACTCAC TATGAGCTTA GCTTTTAAGT TGCTTAAGAA TAGGTAGATC TATGCAAATA ATGATAATTA

TTATTATTAT TTTAAGAGAG GGTCTCACTT TGTCACCCAG GCTGGAGTGC AGTGGTGTGA TTAAGGGTCA CTGCAACCTC CACCTCCCAG GCTCAAATAA ACCTCCCACC TCAGCCTCCC CAGTAGCTGG AACCACAGGC ACGGGCCACC ACGCCTGGCT AATTTTTTGT ATTTTTTGTA GAGATGGGGT TTCATCATGT
TGCCCAGGCT GTTCTTGAAT TCCTCGGCTC AAGCAATCCT CCCACCTTGG

CCTCCCAAAA TGCTGGCATC ACAGGCATGA TGGCATCACT GGCATCACAT ACCATGCCTG GCCTGATTTA TGCAAATTAG ATATGCATTT CAAAATAATC TATTTTATT TGTTGCCTTA TTGGTGGTAC AATCTCAAGT GGAAAAATCT AAGGGTTTTG GTGTTATTTG CTTACTCAAC CAATATTTAT TAGACTCTTA CTAAGCACCA ACATGATCAC ATGCCTGAGC TATGGCTAGC ATAGCGTGTG

AGACAAACTT AATCTCTGTT TTGGTGGAGC ATATAATCTA GTAGATGAAG CCAATGTTGA GCAACATCAC AATACTAACA AATTGAGGAT GCTACGAGAG TGTCTAACAA ATTGAGGATG CTACGAGAGT GTCTAACAAA TTGAGGATGC TATGAGAGTG TGTCATGGAG AGCTGCCTGG AGATTGAGAG AAAGCTTCCT TGAGGGAAGT TACATTTCAG CTGAAACACA CTGCCATCTG CTCGAGGTTT TGTAACTGCA TTCACATCCC GATTCTGACA CTTCACATCC CGATTCTGAC

ACTICACCCA GITACIGICI CAGAGCITGG GICCGCATGI GIAAAACAAG GACAGTATGC ACTTGGCAGG GTTGTGAGAA GGGAAGAGAA CACAAGTAAA GCACCTGTAT CAGGCATACA GTAGGCACTA AGCGTGCGAT GCTTGCTATG ATTATACATC AGTGTAAGCA TCAAGGAAAA GCTGAAGAAA AGTCTGACCA

ACAGCGAAAG ATAAATGCGC AGAGGAGAAA TTTGGCAAAG GCTCCAAATT CAGGGGCAGT CCGTACTCTA CACTTTGTAT GGGGGCTTCA GGTCCTGAGT TCCAGACATT GGAGCAACTA ACCCTTTAAG ATTGCTAAAT ATTGTCTTAA TGAGAAGTTG ATAAAGAATT TTGGGTGGTT GATCTCTTTC CAGCTGCAGT TTAGCGTATG CTGAGGCCAG ATTTTTTCAA GCAAAAGTAA AATACCTGAG

AAACTGCCTG GCCAGAGGAC AATCAGATTT TGGCTGGCTC AAGTGACAAG CAAGTGTTTA TAAGCTAGAT GGGAGAGGAA GGGATGAATA CTCCATTGGA 2200 2250 GGTTTTACTC GAGGGTCAGA GGGATACCCG GCGCCATCAG AATGGGATCT 2300 GGGAGTCGGA AACGCTGGGT TCCCACGAGA GCGCGCAGAA CACGTGCGTC 2350

AGGAAGCCTG GTCCGGGATG CCCAGCGCTG CTCCCCGGGC GCTCCTCCCC

				17	
GGGCGCTCC	T CCCCAGGCC	T CCCGGGCGC	T TGGATCCCG	GCCATCTCCGC	2450
ACCCTTCAA	G TGGGTGTGG	G TGATTTCGT	A AGTGAACGT	G ACCGCCACCG	2500
AGGGGAAAG	C GAGCAAGGA	A GTAGGAGAG	A GCCGGGCAG	CGGGGGGGGG	2550
TTGGATTGG	G AGCAGTGGG	A GGGATGCAG	A AGAGGAGTG	GAGGGATGGA	2600
GGGCGCAGT	CONCCCCTC	A CCACCCON	A AGAGGAGIG	GAAAGGAGAA	2600
A A CCCCCCOOM	G GGAGGGGGIG	M GGAGGCGTA	A CGGGGGGGA	GAAAGGAGAA	2650
AAGGGCGCT	o GGGCTCGGC	G GGAGGAAGT	G CTAGAGCTC	r cgactctccg	2700
CTGCGCGGC	A GCTGGCGGG	G GGAGCAGCC	A GGTGAGCCC	AGATGCTGCT	2750
GCGCTCGAA	CCTGCGCTG	C CGCCGCCGC	GATGCTGCT	CTCCTGGGGC	2800
CGCTGGGTC	CCTCTCCCC	T GGCGCCCTG	CCCGACCTG	GCAAGCACAG	2850
GACGTCGTG	ACCTGGACT	T CTTCACCCAC	GAGCCGCTG	ACCTGGTGAG	2900
CCCCTCCTT	CTGTCCGTC	A CCATTCACO	CARCOCIO	ACGGACCCGC	2900
CCTTCCTCA	r corrections	A CCALLOACGO	CAACCIGGC	ACGGACCCGC	2950
MOCTOCICA.	CCICCIGGG	G TAAGCGCCAC	CCTCCTGGT	CTGTCCCCTT	
TCCTGTCCT	CTGACACCT	A TGTCTGCCCC	C GCCAGCGGC1	CTCCTTCTTT	3050
TGCGCGGAA	A CAACTICAC	A CCGGAACCTC	CCCGCCTGTC	TCTCCCCACC	3100
CCACTTCCCC	CCTCTCATT	C TCCCTCTCCC	TCCCTTACTC	TCAGACCCCA	3150
AACCGCTTTT	TGGGGGGTA	T CATTTAAAA	ATAGATTA	GGGTTACAAG	3200
TGCAGTTCTC	TTCCATGGG	יייני מייני אייני מייני ד	CTCCTCCCN	CIGGGCTCTT	3200
ACTOTA ACTO	TCACCCCAN		GIGGIGGCAI	GGTAATTTCT	3250
AGIGIAACIO	TCACCCGAA	I GIIGIACATI	GTATCTAATA	GGTAATTTCT	3300
CATCCCTCAT	CCCTCTCCC	A CCCTCCCACC	TTTTGGAGTC	TCCAGTGTCT	3350
ACTATTCCAC	TAAGTCCAT	G TGTACACATT	GTTTAGCGCC	CACTCTAAAT	3400
GAGCCTTTTT	GTTTCATTC	A TTCTGTAAGT	GTTGAATAGG	CACCACCTAA	3450
GGTCAGGTAT	AAGTGGAAA'	TTGAAAAAGA	AACTGCCCAC	TTGCCCCAGT	3500
ACTTCCCTAG	CCAAGAGGA	GGAAACCAGG	CACCTCCACC	TGAAGGCCTG	3550
TCACTCCTTC	Victoria Contraction	G CAGTGTAGGA	CAACGAGGACC	TOWNOOFFEE	3550
TONGIGCIIG	ATTIGCTOR	CAGIGIAGGA	CAAGTAAGAT	TGTGCATAGC TTTCTTTTCT AAGGGCGTCA	3600
CIICIGIAII	TAAGACTGT	TTAGGAAGAT	TICTCTTTCT	TTTCTTTCT	3650
TTTTCTTT	TCTTTTCTT	TTTTTTTTA	GGCAGATGAA	AAGGGCGTCA	3700
CAGAACAGGA	ATAAAAATCI	r aaatattcaa	TAAATGAGAC	CTAGGAGACT	3750
ACTGCAGTGA	CTTACAAAGT	г сстаатаааа	AGATGTCTCT	CCAAAATGGG	3800
GCTGCAAAAT	GTGGTGCTGC	CTTATCAGCT	CTAAGTTTTT	TOTTACCTC	3850
AGAAAGAAGG	AACCTGATGC	AGGTTCAGGG	CTCCTCCCC	ATCABTCCAC	3900
GCTGACTCCA	DGDTGGGGA	CTACAGGGAC	PARCOCACCO	ATGAATGCAG	
TOTTATTE A	CCCCCCCCC	CIACAGGGAC	AATCCCAGGT	CTTCTAGGCC	3950
CLIMITING	GCCCTGGGAG	CCTCCAGAGA	TGGCCACATC	TTGACCAGCC	4000
CAGATAGAGG	GAAAGATCAC	CATTATCTCA	CCTCTGTGTC	AAATACCTAG	4050
ATGCTGTCCT	CCCTGAGCCC	ACACTATAGT	TGCCAGCGCT	AATTTAATGG	4100
GTAGTGTACT	GGTTAAGAGA	TGGACAGACC	ATCCTGGCTT	GACTCTCAGC	4150
TCTGGCAAAG	ATGAGTGACT	TGGTTTTTCC	ATATCTCTTG	GCCACACCAA	4200
CCTTGATTTC	TTCAGCTGTA	GAATGGAATT	TCTCAACCTT	GCCTCAAGGA	4250
TTATTGCCCG	ACCATTTCAT	GATATGGTAA	CACCUTCTCA	COCTOROGA	
CATAGTAAGT	CTTTCACCTT	TCAAACGAAT	TOTOTO COMMO	TRACE CAMEC	4300
TCACCAMMO	OTTIONCOIL	ICAAACGAAI	IGITICITIC	TAGGACATGG	4350
IGAGCATITG	GIAGCCATIC	ACCGGTTTTC	TGTTTCTTTG	GATCATAGTT	4400
AACCTCTCCT	TTTCCTTCTG	GCACTACAAT	TTTCTGGTGG	GGAAGAATCC	4450
TTACTTTCTG	CCCTTCCCCT	TAAGGATAGG	AAGCTGATAC	TAGGCAGCAA	4500
CTAGTTGGGG	GATAGGAAGA	TTGTTCCAGA	GAAATGCTGA	ACCATAGGGC	4550
TCCAGATCAC	AGGACCCCAG	TCTTAGCTTG	CTGGGGTGTG	GGGTGGGGGG	4600
GGGCGGTTAC	TGAACATGGG	TATGAAGTAG	ATGTCCATTT	ACTGAAATGT	4650
GAGGACCTGA	CCCCTCTTTCT	ATTCCTCTAC	CCACCATATT	CCCCAACCTC	4700
TCCCCAAGAA	ACCACACATO	GGGGTTCCCC	CCMGCAIAII	CCCCAACCIC	
2000000000	MAGACAGAIG	GGGGTTCCCC	CCTGGAGTAA	CAGGTCCAAA	4750
ACAMAMACA COCA A COMOC	IACAGIGGGA	CTTCCAGGAT	CTGGGCCTGA	TCACCCAGCA	4800
GTCAAGCTCC	CCGCAATTGA	CTAACACCCC	CCTAACACGT	AGAAATTCCA	4850
ATCTGCAATT	TAGTGAGGAT	GATACCTTTA	TTCTTCTTAA	ATACATCTCT	4900
TCATTTCCCA	GAGCACCCTT	TTTTCCCCTC	CTCTGCACCT	TTTTGTTAAA	4950
GACTGGAGTA	TAATGAAATA	CCAAGAGAGC	ATAACATGTG	ATACATAAAA	5000
CTTTTTTTCT	GGTTTACAAA	ACAGTTCATT	CTTGTCCATA	CGTGCTTCTC	5050
TCCAAGGCTG	GCTGCTGTCT	GTTCCAGCCC	CCLLCCCLLAC	GAGAGGCCAT	5100
CTGCCATACC	TECTCCCCAG	ACGCATCGAC	AACCACACCC	ACACECCET.	
CTCCTAACAC	CERTATAGE	ACGCATCGAC .	MAGCACACCC	CTAAGACCTA	5150
COCCOMMONC	TI CI COCOCO	GAGGAACCCC	CICICCICAT	CTAAGACCTA	
GCITCIAAAI	TAGAGTGTGA	GGGTCCATCT	CCCCAGGAGG	GGCACAGGGC	5250
CCAAACAGCC	CAGCCATCTC	AGAAGACAAC	ACTAAGCTTT	GTAGGGGTCC	5300
ACAGTAGAGG	AGAGTAAGAC	GCCTGTTGTT '	TAATTTATTA	CAGTTCCTCA	5350
aaagtgaaga	TGTGTGGGCG	GGATGGCAAG	AGCTGAGCAG	ACGAAAGCTG	5400
AAGGAATAAG	GAAAGAGAGG	AGGACACAAA	CAGCTGACAC	TTCCTCAGTT	5450
CTTGTCATTT	GCCTGGCCCT	GTTCTAAGCA	CTTCTACCT	ATTA ATCCAT	5500
TTAGTCTTGG	CTACAACACT	GTGAGTAACT			
AAAAATTTAAC	AAAGTCACCO	TCACCCACCE !	LY VOIL TIGICA	CCCCCATTT	5550
TCARACTACE.	CONCERNO I GAGGC	TCAGGGAGGT	LAAGTAACTT	GGCCACAGTT	5600
CONCINCIAGA	CICIGATCAC	ATGAGATAAT A	AGTGCCCATA	AAAAGGGAAA	5650
GCAGATTATA '	TTTTTAAAG	GAAAGAGAGT A	AGGATATGGT .	AGAAAAAGAT	5700
TGTTTGGAAA (GGAATTGAGA	GATTGATATA A	ATGAAAAGAA	CATTCACAT	5750
GAGAGTAACA	GTATCAGGGC	CCAAACCTTC A	ATCTAAGGTA	CTTCAAAGAG	5800
GCCTAAGCAA	ACTTAGTCAC	TGGCGTGGTT (TAGTCTCCA '	PGATGGCAAA	5850
TACATTGTGT	ACAGCCCAAC	TCCACACAAA A	מדמממדדם	TANTCATACA	5900
GCAATCTAAA	TTTGAAACA	AAAAATCTTT (
GGGACTTANT	TARCARACCE	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	WHITIGICS :	CITUCCAGA	5950
CACAACTIMAL (ANDROMACUA .	ATCAAAATAC 1	TCCTAAGCC !	MACTGTGTG	6000
CAGAACTCCA	AGAGAGCCC	AGCCCTAAAT C	CAACACTGTC (CAATGGAAAT	6050
ATAATATAAT (FTGGGCCTCA	TATGCAAGGT (ATATGTAAT	TTTAAATTTT	6100
CTAGTAGCCA :	DAAAAATTAT	GTAAAAAGAA A	CAAGTGAAA 1	AATTTTAA	6150

				20	
				G CATGTAATCA	6200
				C AAACCAAGTC	6250
TATTCTATAA	TCTGGCGTG	T ATTATTTAC	A GCACTTCTC	A GACTATATTT	6300
CTTTCTTTCT	TTTTTTTT	C CGAGACAAT	TTGCTCTTG	CACCCAAGCT	6350
				CTCCCGGGTT	6400
CAAGTTATTC	TCCTGCCTC.	A GTCTCCCAA	TAGCTGGGA	TAGAGGCATG	6450
CACCACCACG	CCTGGCTAA	T TGTGTATTT	TAGTAGAGA	AGGGTTTCAC	6500
CATGTTGGCC	AGGCTAATC	T CAAACTCCT	AGCTCAGGTC	ATATGCCCAC	6550
CTCGGCCTCC	CAAAGTGTT	G GGATTACAGO	CGTGAGCCAC	TGCACCCGGC	6600
CTCAGATTAA	CTATATTTC	A AGCGTTCAGT	AGCCACATG	AGCTAGTGCT	6650
ATGGTAGTGG	ACAGTACAG	A TOTGOATTTO	DATTABGACA	CGTATACAAG	6700
CATAGTTCAC	TAATGCACG	מממממממת ב	TATACTCCTC	AGTCGGTGGT	6750
AGAAATCCTA	AATACTGCAG	S AGCAAAAGTG	GTACGAACAG	CAATCTCAGT	6800
				TTTTCCTTCA	6850
		CCAATTCAAT			
				TTTTCACATT	6900
					6950
				CCAGTGTTGG	7000
AAAGAGGIGU .	AGACTCCCC	TGTGCCTATT	GATGGCAGAA	ATATTCACAG	7050
				TGTCTCCTAA	7100
		TCTGATTGAG			7150
		CCTGGAGCCC			7200
		CCTCTACCTC			7250
		TCTTCCCTGT			7300
GCATATTCTC	CCATAGTCCA	GTTCTTTTCC	TGTTCTCCAG	TCTGGCTTCT	7350
GGATGACAGC (CCACTAGTTI	GAACTCCATA	CTGCTATAGT	TCAAGTCCCT	7400
		AAATTACCTC			7450
TTGTAAAATG /					· 7500
TTGAGTGAAA (7550
CTGATGTGCA :					7600
CACATCTGGC 1					7650
TTACTTACTC (CCCTTATTA	ACTGAAGACT	GGCACTGATC	TCACAGTTTC	7700
CTCTCCACTT (CTAGTCTCA	CCATCATCCT	AGATGACTTC	AAGTCACCTA	7750
GATAAACTGT (7800
TACACTCAAG 1	TTGTAACAGA	ACCAGCTTAT	CCAGCTCATG	AAATGTATGC	7850
ATTTCATCTC A					7900
AGCCATGGTG A	GAATATTTA	CCATGGAAAT	TGGCAAATAC	TAAAAAGCAG	7950
AGCACCTTTT 1					8000
CCATCATAAC A					8050
TACTTCTTCC A	TATCTGATT	TGAGCTTCTT	AATTTATCAT	GTGAACCACT	8100
CTTGTAATAA T	AACCCCAAA	TCCCTGTTCC	ATTGTTCTTC	CTGCTAAAAT	8150
ACTAAACCTG G	TTTAGTCCA	ACCATATITT	CTCTCTTTGG	AATCTACAGG	8200
GTGGCCCAAA A	ACCTGGAAA	TGGAAAAATA	TTACTTATTA	ATTTTAATGT	8250
ATATTAATAA G					8300
GTATAGCTGG G					8350
CAGCCACACA G					8400
CCTGGCCCTT A					8450
CTCTACTTTT T					8500
ACATGACCAT A					8550
CCTTTTCCTC C					8600
TTTCCTCCTT T					8650
CCCGTCCCCT C					8700
CCTCTCCTGT A					8750
CTTTCCCCAA G					8800
AGAAACCACC C					
TGCCCTCATG A					8850
ATCATCAATG G					8900
TTTGGTCTTT G					8950
AGTCCTAACC C					9000
					9050
ATTGCAGACG T					9100
GCTTATCTAA TI					9150
GACACGCACA TI					9200
AGTCAAAAAG C					9250
CTGCGCCTAG AC					9300
TCGGCTTTTC A	AACTGTAA	GACAATACAT	TTCTGTTGTT	CAAACCAATT	9350
AGTTTGCAGT AC	TCTGCGAC	TGCAGCCCTA A	ACAAACTAAT	ACAGTCTCTT	9400
GGAGGCATTT GC	CAAGGTTG	ACAATGGAAG (CACTTTCTTA	CCCCTTTAGG	9450
TCTGTCGCCT TT	CTTGTTGG	GGGGTGTTTT (CTAACAATTC (CTCTCCATCT	9500
CTCTCTCTCT AC	TTTGTCTT .	AAACATTGGT (STICTICAGA (CTTCTGACCT	9550
AGGCCTTCTT TT	CACTTCAC .	ATATTCCCCT (GGTGGTCTC A	ACCCACTTCC	9600
AGAAATTACT TA					9650
ACAACTGGCT CT					9700
TTCTGTGGTG TA	AATACTCC	CTCCATGGCC A	AATTCCAAAC :	FGCCAACAGT	9750
TTAACAACTG GC	TCACAAAT '	TTTCTCCAAA 1	TTAACATTT (GCTTTCACA	9800
GGCCAACAAC GI	GGTACAGC (CAACTCCAGC A	ACACCTCTGC :	TTTTGTGTCA	9850
GAGAGAAGTA AC	TTATTTTT	GTACAAAAGG 1	AAAATAAAA	CACCTGCAG	9900

				21	
			T AGAAATAGAA	TAGCTGAAGO	
				G TGGTGGGATT	
				ACCTTTGATO	
				TAGTTTTTGT	
AGGAGACAAA	CAMIAITATI	CTTAATACC	L AAGAAAAIAI	TTCATTGCAA	10150
				TGGGTTTTCA	
				TACATCAGGC	
				CAGCGGTGTG	
CCAAGTACTT	ATCTAGGTAT	CGGGTAGATT	CTGATAAGTC	AGTCAGGTCC	10400
				AGAGAGTAAG	
				AATGCTATGA	
TTTCTACCTC	TARGETCAG	ACTICATIGGAC	AGTGACCCAA	GGTGGTACAG	
				TCCCACTTCC	
CAGTCTCTTG	CCCAGCCGCG	ACTGCTTACA	AATACAGCTA	GAGGAATCTA	10700
AATGAGGTTC	CTCTATCATC	AAACCCAATC	AAAATGCCAA	GGAACAGAAT	10750
CAGTGCCTGG	CTGAAGGCAG	TGGAACAGGG	CCAGCCTGGA	GTGGTTCTCT	10800
CTGAGGAAGT	TCCTCATCTT	GGTTTTAGGG	CCATACCTTG	TGACCTGTGA	10850
GCTAGGGGTT	GCCAGTCCCT	GACATTTCTA	CTGAGGACTC	GCCTGTCTAT	10900
				GGGCGAAGCG	
CCTGATGGAT	GGAAGTATGT	TTTTTGGTGT	TCCATTGGTA	TCTCAAATTC	11000
TACAAAACTT .	AGTGCCCCTT	CTCCTCCCTG	TTCCTCCCCA	TCTTCAGTCT	11050
CCCAGGAGTA	ATCCTTCACT	CCTCCTCAAC	ATCCARCTIC	CAAGGAGCTT TAATCAAATC	
				TTGGGAGGCT	11150 11200
				CCTGGCCAAC	11250
AAGGTGAAAC	CTGTCTCATT	TAAAAAAAGT	TATTTTAAAA	ACTCAAATCT	11300
ATTATTTCTA	CCTCTAAGTG	TGTCTTGAAT	TTATCCATCT	CTCTCCATCT	11350
CTGAGCTGTT 2	ACCTTACCTC	AGTCCATCAC	GTTTTGTCTA	CGTTAACATG	11400
ACCAGAGTCT :	IGTTCTTAGT	CTGGTGAGGT	CACTCCAGCT	GCTTCAGATC	11450
				CCTGGACATG	
				TCCATTCTGT	
GCTCGTCAAT	CICICIGCC	TOTOTTOTA	GGCACCAACC	ACACCCTTCT	
TTTCCCTTCA (CTGGAATGC	TTTCTTCAAT	CCTACCCCAC	TCTCTTTAAT	11650 11700
CTAGATAAGG 1	TTATTCTTT	TTGAATGTCT	AGCAGTGAAA	CCATTTCCCC	11750
TGAAAAACCT 1	CTCTAACCA .	ACCCCCTACC	CTCAGCCCAA	GGTCTAGATT	11800
AGGAGTCCCT (11850
CTTGTTCGTA 1					11900
CTCTGCCTGG T	TCACCATTC .	ATCTCCAGCA	ACTAGCATAA	TGCCTGGCAG	
AGTCAGCCTG C	CAACAAATAT	TTGTTGAATA	AATTAACAGA	TGGCTTTATC	12000
AGGTGTGGTG G	CCCATGCCT	TTTTCACCIA	CACTETCCCA	CCCCCACAGGCC	12050
GGCGGATCAC C	TGAGGTCAG	GAGTTCAAGA	CCAGCCTGGC	CAACATCCTC	12100 12150
AAACCCCATC T					
TGCGTATAGT C	CCAGCTACT A	AGGGAGGCTG	AGGCAAGAGA	ATCGCTTGAA	12250
CCCAGGAGGC A	GAGGTGGCA (GTGAGCCGAG	ATCATGCCAC	TGTACTCCAG	12300
CCTGGATGAC A	GAGACCCTG 1	CTCAAAACA	CACACACACA	CACACACACA	12350
CACACACACA C	ACACACACA (CACACACACC	AAGTTGTATA .	ATTAAAATTA	12400
TAACGTGCTT G	TTATGGAAC A	CTTGTAAAA	TACAGGAAAG	TAATGAAAAA	12450
GTCTACCATC T ATTCTCTCCT G	AGCTCACCA C	CATAATGACC	ATTGCTATCA	TCCTGGCATA	12500
TGAGTACTAT T					12550
CTTGCCATTT T					12600 12650
TGTGCAAATA T					12700
TGAAACTCTG T					12750
TACAATTTAT T					12800
TCCTTACTTA G	TTCAGATTA G	CATTTCCAT	TTATTTAGCC (STGGTTTTGA	12850
GGATGCCATG A	CAGATGCCA T	CCTTCCTAG	AGCTCTTTGG (GCTGTCAGG	12900
TATTTCAGTC A	GGGTGAATT C	GGGTTGATA .	ACATTITAAA 1	ATCTCACTTT	12950
ATTCTGAGGT TO	CCTAGTGTC A	GAGCCCACC	GTATTTTTAG (GACTCCCAA	13000
GTTACAAACA AA	AAATATGGT G	AGGAGGAAT	CACTGAAGTT	TAACACAAG	13050
AGACTTACAT T	TONDART T	COMPCOR	TAGITTATTT (CTAAGCATA	13100
TGAGCACATC T	TAAAAATII I	ACAIAGCAI	TCACATATT 7	AATTAAGCA	13150
ATATTAAGAG G					13250
ATCCCAACAC T					13300
GTGGAGGCTG CA	AATGGCCTG A	GATCACGCC A	ATCGTACTCC A	GCCTGGATG	13350
ATGAGAATGA AA	ATCCTGTCT C	AAAAAAAA	AAAAAAAA	AAAGAAGAA	13400
GAAGAAGTAT TO	GCAATCAG T	GCTCCAGGA A	ATAATTTCCT C	ACTTGAAAT	13450
AAACCTACAT GT	MAGACAAAC T	AATTAGGCC /	ATTCCAAGAG T	TGCTAGCAT	13500
IGGITTAATA TO	JITITCAGA G	CATTCCAGG A	AGCAGTGTG G	CCAGCATTG	13550
CATCETTON					12600
CATGTTTGAT ACTOCTTTTC TT	TTTCAGAAA T	CATCTARRA C	ATTITUTE I	MCCCAGGIC	13600

				22	
TTTTGAGGGA	AGGGATTATA	GATCATTCT	A ATTCCATTT		13700
GTACCATTCT	AAGCACATCA	TACCCACCC	A TTTGGAGCAT	- CINCULIII	13700
363633556	Chromacaton	TAGGCACCC	A IIIGGAGCA	IIIIGGCIIG	13750
ACAGAATATG	CATTIAGAAT	IGITCAAAT	T AGAGGTGTCA	GTGATGGGA	13800
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TTTTCCTTGG	TGGGGAATGG	TGAAGGGAG	G CAGGAGTTAA	GAAGAGGAGA	13900
AGAGATCCTA	AGTCATTTAT	AAACTTCTC	T GGAAAGACAG	GTGTGTGAAG	13950
ACTTTTTAAA	AAGTCATTCA	CCAAATTCT	G TGTGTGTGTG	TOTOTOTO	14000
TTAAATAGAC	Jahar Variation	ACACCACTO	T TAGGTTCACA	0010101011	14000
ITAAATAGAC	111/111111	AGAGCAGIT	1 TAGGITCACA	GCAAAATTGA	14050
ATGCAAGGAC	AGAGATTTCC	CATAAACCC	C CTGCCCACAC	ACATGCATAG	14100
CCTCCCTCAT	TATCAACATC	CCCACCAGAG	G AGGTGTTTGT	TCTAGTTGAT	14150
GAACCTACAC	TGACACATCA	TTATCACCC	A AAGTCCATAG	TTCACGGCAG	14200
GGTTCACTGT	CGGTGTACAT	TCTATGGGT	T TGAGCAAATG	TATAATGACA	14250
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CCCCIGGTAA	CCGCTGATCT	TTTTACTGT	CCATAGTTTC	GGACGATCTA	14400
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ACT TGAAGCC	AGAAGIIIGA	GACCAGCCTG	GGCAACATAG	CAAGACCCCA	14650
			CACACACACA		
CACACACA	CACAAAAACA	AGCTCTTGCC	AGAATTAGAG	CTACAAATTG	14750
CCCTCAGGTT	CCTAGAAGAT	CAGTCCTTCA	ATTAGATTCA	GATTGAGATG	14800
CTTCCTCTTT	TAAACAATGA	TTCCCTTTCT	ATCATGCCCA	ATANGANAC	14850
TAAAAAAAA	TABACAATAC	TCCCTCTAAT	CTCAGCTACC	CACCACCCAC	14000
ANCCACAACT	COURCESTAC	CCCTGIAAI	CICAGCIACC	CAGGAGGCAG	
AAGCAGAAC1	GCTTCAACCC	GGCAAGCAGA	AGTTGCAGTG	AAGTGAGATC	14950
			GCAAGATTCT		15000
CAAAACAATG	TGATTTCCTC	CTCTAAGTCC	TGCACAGGGA	AATGTTAAGA	15050
AATAGGTCCA	CCAGGAAAGA	AGGAAGTAAG	AATGTTTGAC	TAGATTGTCT	15100
TGGAAAAAT	AGTTATACTT	TCTTGCTTGT	CTTCCTAACA	CTTCTCCAAA	15150
GCTTCGTACC	TTGGCCAGAG	COTTGTCTCC	TGCGTACCTG	PCCDMMCCMC	15130
CCACCAACAC	ACACOTTO TO	ADDOCTOR	COLDINCCIG	AGG111GG1G	
GCACCAMGAC .	MONCTICCIA	ATTITUGATO	CCAAGAAGGA	ATCAACCTTT	15250
GAAGAGAGAA	GTTACTGGCA .	ATCTCAAGTC	AACCAGGGTG	AAAATTTTTA	15300
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44444444		CCCCTCCTCC	ATATAACTGA	GGCCTCACAA	15500
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GGTGAACGAG A	ATCTGTGGAC :	TTCTGGGCTC	CAACGTTAGA	TTCTGATTTT	15750
AGCAAGCTTG 1	CAGGGGATT (TGATATTCA	AAGGCTGTGG	CCTTCACCTG	15800
AGAAACCTGC (CTAGGGGGC	יייד א א א מייד איי	TGTCCTGTCT	TOTAL CARCOLO	
CTATCACACA	CANAMOORA		IGICCIGICI	II CAGAAGIG	15850
GGGGGGALGER I	CAAAIGGAA (TIAAATCGT	ATCTTAACAA	TTACTAGGAT	15900
GGGCGCAGTG A	CTCACACCT C	TAATCCCAA	CACTTTGGGA	GGCTGAGGCA	15950
GGAGGATCAC I	TGAGCCCAG C	SAGTTCGGGA	CCAGCCTGGG	CAACATAGAG	16000
AGACGTTGTC I	CTATTTTTT A	ATTAATTAA	AGAGAAAAA :	ATACTGAAAA	16050
TATTGTATAC A	CCACTGAAT 1	DTAATAATA	TGTATATAAT (מדב דב דב דב דב דב דב.	16100
ATTATGAGGA A	מ דייים ביורים ביו	א די א מידער א ידי א מידער א	THE PROPERTY OF		
TATTTTATCC A	CHUNATONNO I	NATIONIAN .	AAMMARCIII .	ICCIICIGII	16150
TATTTTATCC A	GITATGAAG I	ATTTAGAAC	AATTCATCAG	PAAT TGGGGC	16200
TAAATTGACA G	AATAGTAAT (AGAGAAAAT	AGAAAAAGAC A	AGATGGGTTA	16250
TCTTTGAATA C	CAGGTTGGA G	TTGTTTATG	GGTTTGTTTT :	TGTTTTGGG	16300
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TCCCACCTTA G	CCTCCTGAG T	ACCTCCCAC	CACACCTCCA	CTCACCACA	16450
CCCAGCTAAT T	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TTOURCE	CACAGGIGCA I	OI CACCACA	
CCCAGCTAAT T GGCTGATCTC A	**************************************	AUAUAUA	CAGICITICT I	AIGITATUCA	16500
OGCIGATOTE A	AACICCIGC A	CTCAAGTGA	TCCCCCCTGCC 1	TTGGCGTCCC	16550
AAAGTATTGG G	ATTATAGGC A	TAGCCACCA	CACCCAACCT A	GTTTCTATT	16600
TAGACTTGGC C	CTTTCCCAC C	AGTCATTTG	TGTCCAAAAG A	TCTCATAAA	16650
TGTAGACAGG A	AACTGTCCT T	TGCTCATCA	COMPONICA O	つつからからかつつつ	16700
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ATCCTCTCTC	CCCCrccca m	777777	INIGCAMGII C	CICIGAAAC	16750
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TGCAGCCTCT G	CTCCCAGG C	TCAACCCAT	COTOCOGGAL C	COTOTOGCICAC	17000
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CCTGAGCTCA AC	SCTATCCAT C	IGCCTTGGC (CTCCCAAAGA G	CTGGAATTA	17200
CAAGCGTGAG CO	CACTGTGCC TO	GACCAGGGT (GGATTTTTTC A	AGTGCACAT	17250
GTTGTGGTCC CA	GAAGCTCT CI	ATGGTACCA	AATTCCAAGC C	2222222	17200
CAATGGTTCC CA	ACCCATCCT N	יייסייייייי י	**********	A A A TOTA COS	1/300
CACTGCAGAT A	DOTOTAL A	ANNOUNCE A	AIGGLAAGAG G	MAATUACCA	1/350
CACTGCAGAT AC	MOTCCATG TA	MAACAAAT 1	GCTATGGAT T	TTGAAAGTG	17400

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			24	
TTTTGTTTTT TTTTTTA	AG TAGATGTGG	CAGACGTGGT	GGCTCACGCC	21200
TGTAATCCCA GCACTTTC	AG AGGCTGAGG	C AGGTGGATCA	CTTGATGTCA	21250
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AATACAAAAA CTAGCCGC				
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TATAGGGGGT GTATAATA	GA ATTTCGAGCT	' ATGTAAATTC	CAAGTGCATT	21600
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ATCCAGTAGA ATGCTGGG	TO ACAGGACTOT	ACCCAACCTOTT	TCAAACOCAA	22150
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ACTGCAGCCT TGACCTCCC	T GGGCTCAGGT	GATCCTCCCA (CTCAGCCTC	23350
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AGTGCAGTGG CGCCATCTCG	GUICACIGUA A	GUICCACCT C	LCAGGTTCA	24900

25	
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CCACCACGCC CGGCTAATTT TTTTGTATTT TTAGTAGAGA CGGGGTTTC	A 25000
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ATTAGATATG GAATATAGTC TAGTTCCCAG ATATCCATAT CCATTGGTTT ATTACCCTCA TTATTAACTT CAAATTGTTT AATAGACCCT CATATCTCAG	26050
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CTATGAGTTT TACTTTACTT ATTTATTTTA TTTTTTGAGA CAGACGCTTG	26150 26200
CTCTGTCACT CAGGCTGGAG TGCGGTTGCG TGATCATGGC TCACTATGGC	26250
CTCGACCTTC TGGGCTCAAG TGATCCTCTC CCTCAGCCTC CCAAGCTGAG	26300
ACTACAGGCA TGCACCACCA CATCTAGCTA ATTTTTTTT TTCCCCATGG	26350
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CATAGCGCCA GACCTGGTTT TACTTTTCTT GACTTTGAAT TACAAGTTTT	26450
TGTAATTTGG AAAATGTTTT GTTGCTTTTA AATACTGCTG TATGTTTGCT	26500 26550
TTTAAATACA ACATTTCTCG ATATATATTT TGAGAATTGC TGTCTTTCAG	26600
AACCTAACAG TTTCCTTAAG AAGGCTGATA TTTTCATCAA TGGGTCGCAG	26650
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CAAAAATGCA AAACTCTATG GTCCTGATGT TGGTCAGCCT CGAAGAAAGA CGGCTAAGAT GCTGAAGAGG TAGGAACTAG AGGATGCAGA ATCACTTTAC	26750
TITTCTTCTT TTTCCTTTTG AGACAGAGTC TCACTCTGTC AGCCAGACTG	26800
GAGTGCAGTG GTACAATCAT GGCTCACTGC AACTTCGACC TCCCAGGCTC	26850 26900
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TTACCAAAAA AAGGAAATTT TGACGGGTTC AGAATATCAA GGGATCTGAG	27250
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CIMIGRIAIC	COTACAACOT	TTCCTCTGTC	CCCTGTGTTT CAGGTTTTTA	CCTGTAAACT	34300
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COMPAGENCE	CCCAGGATAA	CAGTGATACT	CTTTGTAGGA	FAACTATTTG	35150
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GCCTTTTACC	TTCCTATGTC	TGAAAATGGA	TAGTCTGGCT G	CCTCTTAA	36100
CAACCCAGCT	GGCAGAGCTG	TGAGGATCTC	AGTGTGCTCT A	GCCCAGACA	36150

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TACTTGCGGT TACCCTATCC TTTTTCTAAC AAGCAAGTGG ATAAATACCT 36500 TCTAAGACCT TTGGGACCTC ATGGATTACT TTCCAAGTAA GTAATTTTCC 36550 TTGTTCATTC CAAACTTTCA ATAAATTTAT TGGTGTTTAT CAGAATAGAG 36600 AGTTTGGACA GGGAGCAAAA GACAAAGTCA ACTATATCAA GTTCTAATAA 36650 TTCTTAATAT TCAGGAAATT TATGTATGAA TACTTACTAA TATGAGTATA ACTCATCCTA AGAGTCTAAA GCAAAAGGAT GTGAACACAA ACTAGCAGTT 36750 ATCTTAGAGA ATAAGTTTGC ATTTCAAAAT AACTTGACAT ATCAAGATCC 36800 ACTCAACGCA TTTAAATTAT TTACTCTAAA AAGACATAAT TCTTGGTAAC 36850 ACATTCACTA AAGCAAAATA TACCTTTATA TAATTGCTAT CAAAGGTATG 36900 TGGGTTGGTA TAAAATATCA TACCATGTGA GATCAGTGTG ATTCCTTTAC 36950 AGCATTAATT TTTATTGGTT AGAGTAAGAA AAAGAATAGC TAGAGTATAT 37000 TTCTTAAGTA GATTCTCATA CACTTTGGTT TCAAAAACCA ATTATTGACT 37050 ACATCTTATA AAAGCCTGTA TTCAATGGAG TGCCAAAAAA TGACTATGAG 37100 TCTTAAAGAG TTAGGCATAT AAATATTTTA AGGTTTCTGT TCAATGTATG 37150 TTGGAAGGAG TTCCTTTCTC ATGACTATTC TCATATTGGA GCATAAAAAG 37200 AGTTTACAGG CTTGGCGCAG TGGCTCATGC CTGTAATCCC AATACTTTGG 37250 GAAGCTGAAG CAGGCAGATC ACTTCAGCCC AGGAGTTTGA GACCAGCCTG 37300 GGCAATATGG CAAAACTCTC TCTACAAAAT ATACCAAAAT TAGCCAGGCG 37350 TGGTGGTGCA TGCCTGTAGT CCCAGCTACT TGGGAAGCTG AGGTGGGAGG 37400 ATTGCTTGAG CCCAGGGGGG TCATGGCTGC AGTGAGCTGT GATGGTGCCT 37450 CTGTCACCCA GCCTGGGTGA CAGAGTGAGA CCCTGTCTCA AAAAAATAAA 37500 TAAATAAAAA TTAAGAGTTT ACAAAATTCT CACCATCTCC TCCCATCTTT 37550 GCAAATGCCA CATAAGTGAT GTGTTCCAGG ACTATTAGCC TCGGAACCTG 37600 AGGCAGTACA GTAAGCACGC TTTCTCCAAA GTCCTGTCCC CCACAGACAA ACATTATTTA CACTGGGTAC TGCTCTTTTA TTTTTTCCCC TCTATGCTTT 37700 ATTITACTAT AACTATAATC ATATAACATG TAATAGGAAA AAGGCAGGGT 37750 CGGGGGAGAG ATCCAGAAGT CTTCCCAAGA GCCTTTCCAA CATAGCCTCT 37800 GTAGACATTT TTTCTTTCTT CTTTTTTTTT TTTTTTTTT TTCTGAGACA 37850 GAGTCTCACT CTGTTGTCCA GGCTAGAGTG CAGTGGCGTG ATCTAGGCTC 37900 ACTGCAACCT CCGCCTCCTG GGTTCAAGCA ATTCTCCCAC CTCAGCCTCC 37950 CTAGTAGCTG GGATTAGAGG CATGCATCAC CACGCCTGGC TAATTTTTGT 38000 ATTTTTAGTA GAGATGAGGT TTCACCATGT GGGCCAGGCT GGTCTTGAAC 38050 TCCTGACCTC AAGTGATCCA CCTGCCTTAG CCTCCCAAAG TGCTAGGATT 38100 ACACGAGTGA GCCACCGTGC CCTGCCCCTA TTACATTCTG ATCACACATT 38150 TCATGTTTTA TAATTGGAAA ACTGGTGAAA TTATAGACAA TGTTTTGTTC 38200 CCCTAAATTC TCTTTGATGA GTATATATTA CTTACACTCT TCTGTCTTTA 38250 AAATTTTGCA AAATAGTATC CTAGATAAGT TTATGAGTGC ACAGTCTGTA 38300 CGCTTACTCA TATTAATGAC CTCGGAGAGT TAAACAACAG TCACCTTTAA 38350 AAATTATTAC TATCATTATC ATTATTTTTG AGGCGGGGGT CTCATTCTGT 38400 CTCCCAGGCT GGAGAGTAGT GGTGCGGTCA CAGCTCACTG CAGCCACCGC 38450 TACCTGGGCT CAAGTGATCC TTCCTCCTCA GCCTTCTGAG TAGCTGAGAC 38500 CACAGGCTTA TGCTACCACA CCTGGCTAAT TTTTTAACTT TTTGTAGAGA 38550 CGATGTCTCA TTATGTTGCC CAGGCTGGTC TCAAACTCCT AAGCTCAAGT 38600 GATCTTCCTC AGCCTCCCAA AGTGCTGGGA TTACAGGCAT GAAAAACTGC 38650 ACCCAGCCCT AAAAATTATT AGGGTCCTGC ATAGTAAGAC TTTAATAAAT 38700 ATTTAAATGA ACATCTGGTT TTTTTAAAAA AAAAATAGAG ACAAGGTCTC 38750 ACTATATTGC CCAAGCTGGT CTCGAACTCC TGGACTCACG CAATCCTGCT 38800 GCCTTAGCCG CCCAAAGTGC TGGGATTACA GGCATGACCC ACCTCATCTG 38850 GGCTGAGTGA ACATATTTTT AACATAAAGG CCGTATTTTA TATTTATCTC 38900 ATACATTTTG CCCAGCATCC CCATTTCCGC CGAATCTGTT GCTTGCTAAT 38950 TCCTTCCAGC TTCATTTCAT CTGAAATTTG ACAAACATCT TCTATTTCTT 39000 TGTCGTCATG TTATTGACTT CAGAATATAA AATAAAACAC TATACCCAAA 39050 TTAAACCCCA CCCTCATTGC CCAGCCTGAT GTGAAAATAA TCAGCATACA 39100 TTAAGCTTAC CCTTGATATA TGTGTAGCAT CTTTTAGATA AATATACAGC 39150 TGATTAAGCA ATATAGCCTG ATGGTATAAT ATCTTGCCCA TGTACCTCAT 39200 CTTATCTCCA GCAGGATTAA TTCACAGTGA TCAGATTTAC CTTTAAACTT TGTAGCAAAA TATCCTCTCC AAAAGCATAT CTAAAACTTT TGTGTGTACT 39250 39300 CTTGCAAGTT TCTTAATTTC ATGCAGAACA GGCTCTTACC ACTGTTAGCT 39350 GGAGATATTT TCAAGACCTA TTTTTGTTTG TGGTTTCCTG ATGATGGTCA 39400 TGGCATTTCC CCCTTCACTC CATCTAAAAA TTGAGGTGAT ACAGGCTTTT 39450 AAACAAAACC AACTCATATA GACTGAGTAC AACTGCAATG CAGGCATGCT 39500 AACCTCTGCT ACAATCATGG GCGTGCTATT GATATGTCTT AAGTTACAGA 39550 ACACAGGGCT GAGCGTCTCA TTAGGTCAAA ATGTAAACCA GTTTTTCTGC 39600 TCACTGATGC TTAATGAGGA CAGGGTGTGA GAGATTTCTT TAAGGAAAAC AAATATATAA TAATGCTACA TGGAAAAATA TCTAACATTA GAGAATTAAG 39700 TAAATAAACT AATATACTCA CACCATGGAA TCTTGTGCAG ACATTAAAAT 39750 TATGTAGTGG ATGGATGTTT AATGGTGTGA GAAAAAGTTA GGATGTGCTG 39800 GGGTGGGGG AAGAATCAAG TTTTAAGAAA ATACAGTATA CCCATACTTA 39850 AGTAAAAAA AAAAAAAGG TATGTACAGT CATGTGTTGC TTAATGATGG

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GGATACATTC	CGAGAAATG	T GTCGATAGG	T GATTTCATC	C TTGTGTGAAC	39950
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AGATGAAAGA	ATGAATATA(C ATCAAAATA	T TTAAAATGG	T TATAATGACT	40350
				A GATAATACTT	40400
TTATAGTGTT	TACTATATA	A AAGACACTG	T TATAAGTGT	T CTACATACTT	40450
TATACGTTCT	COTTANAIGA	TATAAATATA	A ACTUIGACA P COPPOPERA	G TAACTAATCT G ACAGAATCTT	40500
				C TGCAACCTCC	40550 40600
				G AGTAGCTGGG	40650
ACTACAGGCA	CACACCACC	A TGCCCGGCT	A ATTTTTGTA	T TTTTGGGTAG	40700
AGATGGAGTT '	TTGCCATGTT	GGCCAGGCT	3 ATCTTGAAC	T CCTGGCCTCA	40750
AGTGATCTGC (CTGCCTCAGO	CTCCCAAAGT	GCTGGGATT	A CAGGTGTGAA	40800
ACTITICITIES (CGGCCTAATC	COTOTTO	TCAATATTT	A AAGAGTGCTA F ATAAGCATCT	40850
TATTTAGAAT	TATGAAAAT	TCARTIGAC	TACACCCCA	TAAAGCATCT	40900 40950
CTTCATAAGC T	CTTGCCTAT	ATTGATTCG	TCCTGTGAA	T ATGCATTAAT	41000
TTGATTTAAA T	FAATAAGTAT	GTATAAGAAA	TAACACTTT	CCTTAATTTT	41050
TAAGAACGTT (CAACAGTTTT	TAATTTGAAT	TCCAATAGT	AAATACATAG	41100
				TTTCACCACA	41150
GCATTCTACC A	AAAATTTCTT	CCCACCTO	AGAAAATGA	TGCATACCTC	41200
AATTTCATTG (CTACCATTY	DOCOTABATT	GGGCATAGT	CONTROL	41250 41300
CTATATATCT A	ATTTTCTTTA	CATAAAAAA	GTTTCAATT	TTGGCCATTA	41350
AATAAAATAG C					41400
CCACCATCAT A					41450
ATGGGCCAGA C	AGTAAGTAT	TTCTGGCTTT	GGAGTCCATA	TGGTCTCTAT	41500
CATAACTACT C GCCTAAGTGA I	ATCTCTGCC	ATTGTAGCTT	AAAGATTATC	TAGGTCAAAT	41550
TAAAAAAAA	TATTTGGTC	TADADADAGAT	TAIMIAMIAI	TTGTAGCACC	41600 41650
ATGGGTGGGG C	ATGCACCAC	TTGGTTAACT	CGGTGTATCT	TTCTCCTTTG	41700
CAGATCTGTC C	CAACTCAATG	GTCTAACTCT	AAAGATGGTG	GATGATCAAA	41750
CCTTGCCACC T					41800
TTGCCAGCTT T					41850
TGCTTGCATC T GTATACTAAG A	GAAAATAAA	CTCAACTAGT	CCTGACACTG	AATTTTTCAA	41900
TGCAAAGCAA C					41950 42000
AGATTTAGCA C	AGTATTTTG	ATCTCGCTAG	GTAGAACACT	GCTAATAATA	42050
ATAGCTAATA A	TACCTTGTT	CCAAATACTG	CTTAGCATTT	TGCATGTTTT	42100
ACTITIATET A					42150
TTGAGACAGA A					42200
CTTGGCTCAC T GCTGGGATTA T.	AGGCGTGTG	AGCAATTCTC	CIGCCTCAGC	TTCCTGAGTA	42250
TGTAGAGATG G	AGTTTCGCC	ATATTGGCCA	AGCTGGTCTC	GAACTCCTCT	42300 42350
CCTCGAACTC C	TGTCCTCAA	GTGATCCACC	CGCCTCAGCC	TCTCAAAGTG	42400
CTGGGATTAC A	GGTGTGAGC	CACCACACCC	AGCAGTGTTT	TATTTTTGAG	42450
ACAGGGTATC A					42500
ATCACTGCAG C TCCCAAGTAG C					42550
AAAAAATTTT T					42600 42650
TGAACTCCTG G	ACTCAATTG	ATCCTCCCAC	CTTGGCCTTC	CAGGTGCTGG	42700
GATTTCTTTG GO					42750
CTCTGTGCAG TO	GTTGCTAGT	CAGCGAAAGA	CTATAATACC	TGTGGGGACA	42800
SCGATTAGCC AC	CACAACCA	GTCTTTATTT	AAAGTTATTA	AAAATGGCTG	42850
GGCGCAGTGG CT	CACACCTG	TAATCCTAGC	ACTTTGGGAG	GCCGAGGCAG	42900
ATGGATCACC TO AACCCCATCT CT	PACGIGAGG	AATTIGAGAC	TAGCTCGGTC	AACATGGTGA	42950 43000
STCCCAGCTA CT					43050
CAGAGGTTG CA	AGTGAGCCG	AGATTGTGCC	ACTGCACTCC	AGCCTGGGTG	43100
ACAGAGAGAG AT	TCCATCTC .	AAAAAAACAA	GTTATTAAAA	ATGTATATGA	43150
ATGCTCCTAA TA	TGGTCAGG	AAGCAAGGAA	GCGAAGGATA	TATTATGAGT	43200
TTTAAGAAGG TO	CTTAGCTG	TATATTTATC	TTTCAAAATG	TATTAGAAGA	43250
ALLINGAATT CI	COTTACCOT A	TGTGCCATCT	CTACAGGCAC	CCATCAGAAA AACTATCTAT	43300
TTGCACCTTA AA	AGACAGCT	AGATTTTGCT	GATTYTTCTTC	TTTCCCTTT	43350
TTTGTCAGC AA	TAATATGT	GAGAGGACAG	ATTGTTAGAT	ATGATAGTAT	43450
VAAAAATGGT TA	ATGACAAT :	TCAGAGGCGA	GGAGATTCTG	TAAACTTAAA	43500
TTACTATAA AT	GAAATTGA !	TTTGTCAAGA	GGATAAATTT	TAGAAAACAC	43550
CAATACCTT AT					43600
CITGITITCA GT	TGGGAAGC :	TTTGGCTGC	AAGTAACAGA	AACTCCTAAT	43650

30 TCAAATGGCT TAAGCAATAA GGAAATGTAT ATTCCCACAT AACTAGACGT TCAAACAGGC CAGGCTCCAG CACTTCAGTA CGTCACCAGG GATCTGGGTT CTTCCCAGCT CTCTGCTCTG CCATCTTTAG CGCTGGCTTC ATTCTCAGAC 43800 TCTGGTAGCA TGATGGCTGT AGCTGTTTCA TGGGCCCCTT CAAACCTCAT 43850 AGCAACCAGA GGAAGAAAAT GAGCCATTTT TTGAGTCTCC TTCATAGACT 43900 TGAATAACTC TTTTTCAGAG CTTCTCACAG CAAACCTCTC CTCATGTCTC 43950 CTCATGTCTT ATTGTTCAGA AATGGGTAAT GTGGCCATTT CACCAGTCAC 44000 TGCCAACAAC AACGAGGTTC CTATAATTGT CTCTGAGTAA CCCTTTGGAA TGGAGAGGGT GTTGGTCAGT CTACAAACTG AACACTGCAG TTCTGCGCTT 44050 44100 TTTACCAGTG AAAAAATGTA ATTATTTTCC CCTCTTAAGG ATTAATATTC 44150 TTCAAATGTA TGCCTGTTAT GGATATAGTA TCTTTAAAAT TTTTTTTTT 44200 AATAGCTTTA GGGGTACACA CTTTTTGCTT ACAGGGGTGA ATTGTGTAGT 44250 GGTGAAGACT CGGCTTTTAA TGTACTTGTC ACCTGAGTGA TGTACATTGT 44300 ACCCAATAGG TAATTTTCA TCCATTACCC TCCTTCCGCC CTCTTCCCTT 44350 CTGAGTCTCC AACATCCCTT ATACCACTGT GTATGTTCTT GTGTACCTAC 44400 AGCTAAGCTT CCACTTATAA GTGAGAACAT GCAGTATTTG GTTTTCCATT 44450 CCTGAGTTAC TTCCCTTAGG ATAACAGCCC CCAGTTCCGT CCAAGTTGCT 44500 GCAAAATACA TTATTCTTCT TTATGGCTGA GTAATAGTCC ATGGTACATA 44550 TATACCACAT TTTCTTTATC CACTTATCAG TTGATGGACA CTTAGGTTAA 44600 TTCCATTCAA TTTCATTCAA TTTAAGTATA TTTGTAAGGA GCTAAAGCTG 44650 AAAATTAAAT TTTAGATCTT TCAATACTCT TAAATTTTAT ATGTAAGTGG 44700 TTTTTATATT TTCACATTTG AAATAAAGTA ATTTTTATAA CCTTGATATT 44750 GTATGACTAT TCTTTTAGTA ATGTAAAGCC TACAGACTCC TACATTTGGA 44800 ACCACTAGTG TGTTGTTTCA CCCCTTGTTA TACTATCAGG ATCCTCGA 44848 (2) INFORMATION FOR SEQ ID NO:43: SEQUENCE CHARACTERISTICS: (A) LENGTH: 2396 (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear SEQUENCE DESCRIPTION: SEQ ID NO:43 (xi) TTTCTAGTTG CTTTTAGCCA ATGTCGGATC AGGTTTTTCA AGCGACAAAG 50 AGATACTGAG ATCCTGGGCA GAGGACATCC TAGCTCGGTC AGATTTGGGC 100 AGGCTCAAGT GACCAGTGTC TTAAGGCAGA AGGGAGTCGG GGTAGGGTCT 150 GGCTGAACCC TCAACCGGGG CTTTTAACTC AGGGTCTAGT CCTGGCGCCA .200 AATGGATGGG ACCTAGAAAA GGTGACAGAG TGCGCAGGAC ACCAGGAAGC 250 TGGTCCCACC CCTGCGCGGC TCCCGGGCGC TCCCTCCCCA GGCCTCCGAG 300 GATCTTGGAT TCTGGCCACC TCCGCACCCT TTGGATGGGT GTGGATGATT 350 TCAAAAGTGG ACGTGACCGC GGCGGAGGGG AAAGCCAGCA CGGAAATGAA 400 AGAGAGCGAG GAGGGGAGGG CGGGGAGGGG AGGGCGCTAG GGAGGGACTC 450 CCGGGAGGGG TGGGAGGGAT GGAGCGCTGT GGGAGGGTAC TGAGTCCTGG CGCCAGAGGC GAAGCAGGAC CGGTTGCAGG GGGCTTGAGC CAGCGCGCCG 550 GCTGCCCCAG CTCTCCCGGC AGCGGGCGGT CCAGCCAGGT GGGATGCTGA 600 GGCTGCTGCT GCTGTGGCTC TGGGGGCCGC TCGGTGCCCT GGCCCAGGGC 650 GCCCCGCGG GGACCGCGC GACCGACGAC GTGGTAGACT TGGAGTTTTA 700 CACCAAGCGG CCGCTCCGAA GCGTGAGTCC CTCGTTCCTG TCCATCACCA 750 TCGACGCCAG CCTGGCCACC GACCCGCGCT TCCTCACCTT CCTGGGCTCT CCAAGGCTCC GTGCTCTGGC TAGAGGCTTA TCTCCTGCAT ACTTGAGATT 850 TGGCGGCACA AAGACTGACT TCCTTATTTT TGATCCGGAC AAGGAACCGA 900

CTTCCGAAGA AAGAAGTTAC TGGAAATCTC AAGTCAACCA TGATATTTGC AGGTCTGAGC CGGTCTCTGC TGCGGTGTTG AGGAAACTCC AGGTGGAATG

GCCCTTCCAG GAGCTGTTGC TGCTCCGAGA GCAGTACCAA AAGGAGTTCA

AGAACAGCAC CTACTCAAGA AGCTCAGTGG ACATGCTCTA CAGTTTTGCC
AAGTGCTCGG GGTTAGACCT GATCTTTGGT CTAAATGCGT TACTACGAAC

CCCAGACTTA CGGTGGAACA GCTCCAACGC CCAGCTTCTC CTTGACTACT

GCTCTTCCAA GGGTTATAAC ATCTCCTGGG AACTGGGCAA TGAGCCCAAC

AGTTTCTGGA AGAAAGCTCA CATTCTCATC GATGGGTTGC AGTTAGGAGA

AGACTTTGTG GAGTTGCATA AACTTCTACA AAGGTCAGCT TTCCAAAATG

CAAAACTCTA TGGTCCTGAC ATCGGTCAGC CTCGAGGGAA GACAGTTAAA

CTGCTGAGGA GTTTCCTGAA GGCTGGCGGA GAAGTGATCG ACTCTCTTAC

ATGGCATCAC TATTACTTGA ATGGACGCAT CGCTACCAAA GAAGATTTTC

1000

1050

1150

1200

1250

1300

1350

1400

				•	
				AAAAATTCTG	1550
AAGGTCACTA	AAGAGATCAC	ACCTGGCAAG	AAGGTCTGGT	TGGGAGAGAC	1600
			GCTGTCCAAC		1650
CTGGCTTTAT	GTGGCTGGAT	AAATTGGGCC	TGTCAGCCCA	GATGGGCATA	1700
GAAGTCGTGA	TGAGGCAGGT	GTTCTTCGGA	GCAGGCAACT	ACCACTTAGT	1750
GGATGAAAAC	TTTGAGCCTT	TACCTGATTA	CTGGCTCTCT	CTTCTGTTCA	1800
AGAAACTGGT	AGGTCCCAGG	GTGTTACTGT	CAAGAGTGAA	AGGCCCAGAC	1850
AGGAGCAAAC	TCCGAGTGTA	TCTCCACTGC	ACTAACGTCT	ATCACCCACG	1900
ATATCAGGAA	GGAGATCTAA	CTCTGTATGT	CCTGAACCTC	CATAATGTCA	1950
CCAAGCACTT	GAAGGTACCG	CCTCCGTTGT	TCAGGAAACC	AGTGGATACG	2000
TACCTTCTGA	AGCCTTCGGG	GCCGGATGGA	TTACTTTCCA	AATCTGTCCA	2050
ACTGAACGGT	CAAATTCTGA	AGATGGTGGA	TGAGCAGACC	CTGCCAGCTT	2100
TGACAGAAAA	ACCTCTCCCC	GCAGGAAGTG	CACTAAGCCT	GCCTGCCTTT	2150
TCCTATGGTT	TTTTTGTCAT	AAGAAATGCC	AAAATCGCTG	CTTGTATATG	2200
AAAATAAAAG	GCATACGGTA	CCCCTGAGAC	AAAAGCCGAG	GGGGGTGTTA	2250
TTCATAAAAC	AAAACCCTAG	TTTAGGAGGC	CACCTCCTTG	CCGAGTTCCA	2300
GAGCTTCGGG	AGGGTGGGGT	ACACTTCAGT	ATTACATTCA	GTGTGGTGTT	2350
CTCTCTAAGA	AGAATACTGC	AGGTGGTGAC	AGTTAATAGC	ACTGTG	2396

(2) INFORMATION FOR SEQ ID NO:44:

- SEQUENCE CHARACTERISTICS:
 - (A) LENGTH:
 - (B)
 - TYPE: amino acid

535

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44

Met Leu Arg Leu Leu Leu Trp Leu Trp Gly Pro Leu Gly Ala 5 10 Leu Ala Gln Gly Ala Pro Ala Gly Thr Ala Pro Thr Asp Asp Val

20 25 Val Asp Leu Glu Phe Tyr Thr Lys Arg Pro Leu Arg Ser Val Ser

35 40 Pro Ser Phe Leu Ser Ile Thr Ile Asp Ala Ser Leu Ala Thr Asp

50 55

Pro Arg Phe Leu Thr Phe Leu Gly Ser Pro Arg Leu Arg Ala Leu 65 70

Ala Arg Gly Leu Ser Pro Ala Tyr Leu Arg Phe Gly Gly Thr Lys 80 85

Thr Asp Phe Leu Ile Phe Asp Pro Asp Lys Glu Pro Thr Ser Glu 95 100 105

Glu Arg Ser Tyr Trp Lys Ser Gln Val Asn His Asp Ile Cys Arg 110 115

Ser Glu Pro Val Ser Ala Ala Val Leu Arg Lys Leu Gln Val Glu

125 130 135 Trp Pro Phe Gln Glu Leu Leu Leu Arg Glu Gln Tyr Gln Lys

140 145 150 Glu Phe Lys Asn Ser Thr Tyr Ser Arg Ser Ser Val Asp Met Leu

155 160 Tyr Ser Phe Ala Lys Cys Ser Gly Leu Asp Leu Ile Phe Gly Leu

170 175 Asn Ala Leu Leu Arg Thr Pro Asp Leu Arg Trp Asn Ser Ser Asn

185 190

Ala Gln Leu Leu Asp Tyr Cys Ser Ser Lys Gly Tyr Asn Ile 200 205 210

Ser Trp Glu Leu Gly Asn Glu Pro Asn Ser Phe Trp Lys Lys Ala 215 220

His Ile Leu Ile Asp Gly Leu Gln Leu Gly Glu Asp Phe Val Glu

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32
                230
                                   235
 Leu His Lys Leu Cen Arg Ser Ala Phe Gln Asn Ala Lys Leu
                245
                                  250
 Tyr Gly Pro Asp Ile Gly Gln Pro Arg Gly Lys Thr Val Lys Leu
               . 260
                                   265
 Leu Arg Ser Phe Leu Lys Ala Gly Gly Glu Val Ile Asp Ser Leu
               275
                                  280
 Thr Trp His His Tyr Tyr Leu Asn Gly Arg Ile Ala Thr Lys Glu
                290
                                  295
                                                      300
 Asp Phe Leu Ser Ser Asp Ala Leu Asp Thr Phe Ile Leu Ser Val
               305
                                  310
                                                      315
 Gln Lys Ile Leu Lys Val Thr Lys Glu Ile Thr Pro Gly Lys Lys
                320
                                   325
                                                      330
Val Trp Leu Gly Glu Thr Ser Ser Ala Tyr Gly Gly Gly Ala Pro
               335
                                  340
Leu Leu Ser Asn Thr Phe Ala Ala Gly Phe Met Trp Leu Asp Lys
               350
                                   355
Leu Gly Leu Ser Ala Gln Met Gly Ile Glu Val Val Met Arg Gln
               365
                                  370
Val Phe Phe Gly Ala Gly Asn Tyr His Leu Val Asp Glu Asn Phe
               380
                                  385
                                                     390
Glu Pro Leu Pro Asp Tyr Trp Leu Ser Leu Leu Phe Lys Lys Leu
               395
                                 400
                                                     405
Val Gly Pro Arg Val Leu Leu Ser Arg Val Lys Gly Pro Asp Arg
               410
                                  415
Ser Lys Leu Arg Val Tyr Leu His Cys Thr Asn Val Tyr His Pro
               425
                                  430
                                                    435
Arg Tyr Gln Glu Gly Asp Leu Thr Leu Tyr Val Leu Asn Leu His
               440
                                 445
                                                    450
Asn Val Thr Lys His Leu Lys Val Pro Pro Pro Leu Phe Arg Lys
               455
                                 460
Pro Val Asp Thr Tyr Leu Leu Lys Pro Ser Gly Pro Asp Gly Leu
               470
                                 475
                                                     480
Leu Ser Lys Ser Val Gln Leu Asn Gly Gln Ile Leu Lys Met Val
               485
                                 490
                                                     495
Asp Glu Gln Thr Leu Pro Ala Leu Thr Glu Lys Pro Leu Pro Ala
              500
                                505
Gly Ser Ala Leu Ser Leu Pro Ala Phe Ser Tyr Gly Phe Phe Val
               515
                                 520
Ile Arg Asn Ala Lys Ile Ala Ala Cys Ile
```

INFORMATION FOR SEQ ID NO:45: (2)

SEQUENCE CHARACTERISTICS:

(A) LENGTH:

TYPE:

(B)

nucleic acid STRANDEDNESS: double

(C) linear

(D) TOPOLOGY:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45

TGC TTT TAG CCA ATG TCG GAT CAG GTT TTT CAA GCG ACA AAG AGA TAC TGA GAT CCT GGG CAG AGG ACA TCC TAG CTC GGT CAG ATT TGG 98 GCA GGC TCA AGT GAC CAG TGT CTT AAG GCA GAA GGG AGT CGG GGT 143 AGG GTC TGG CTG AAC CCT CAA CCG GGG CTT TTA ACT CAG GGT CTA 188 GTC CTG GCG CCA AAT GGA TGG GAC CTA GAA AAG GTG ACA GAG TGC 233 GCA GGA CAC CAG GAA GCT GGT CCC ACC CCT GCG CGG CTC CCG GGC

GC'	r cc	C TC	c cc	A GG	CTC	CG#	GG/	TC	r TG	TA E	r cr	G GC	CAC	TCC	323
GC	A CC	TT	r GG	A TG	GT(TGC	aTC	AT:	r TC	A AA	A GT	GA	GTO	G ACC	368
GC	GCC	GA	G GG(G AA	A GCC	AGO	: ACC	GA	A ATO	AA E	A GAG	G AG	C GAG	GAG	413
GGG	G AG	GC	G GGG	G AGO	GG#	GGG	CGC	TAC	GG/	A GG	3 AC	r cc	C GGC	AGG	458
GG	r GGC	AG(GA'	r .GG/	A GCG	CTC	TGG	GAC	GGT	r AC	r GAG	TC	TG	GC CGC	503
CAC	age	G CG	A AGO	C AGO	ACC	GGT	TGC	: AGC	GGG	CT	r GAC	CC	A GCC	CGC	548
CGC	CTC	CC	CAC	CTC	TCC	: CGG	CAG	CGG	GCC	GTO	CAC	CC3	k GG7	GGG	593
														GCC	638
														Ala	
				5			•		10					15	
CTG	GCC	CAC	GGG	GCC	ccc	GCG	GGG	ACC	: GCG	ccc	ACC	GAC	GAC	GTG	683
														Val	
			-	20			•		25			•		30	
GTA	GAC	TTO	GAG	TTI	TAC	ACC	AAG	CGG	CCG	CTC	CGA	AGC	GTG	AGT	728
														Ser	
	-			35				3	40					45	
														•••	
ccc	TCG	TTC	CTG	TCC	ATC	ACC	ATC	GAC	GCC	AGC	CTG	GCC	ACC	GAC	773
											Leu				
				50					55					60	
														•	
CCG	CGC	TTC	CTC	ACC	TTC	CTG	GGC	TCT	CCA	AGG	CTC	CGT	GCT	CTG	818
											Leu				
				65			,		70	5				75	
GCT	AGA	GGC	TTA	TCT	CCT	GCA	TAC	TTG	AGA	TTT	GGC	GGC	ACA	AAG	863
											Gly				
	J	-		80			-,-		85		,	,		90	
ACT	GAC	TTC	CTT	ATT	TTT	GAT	CCG	GAC	AAG	GAA	CCG	ACT	TCC	GAA	908
											Pro				
	-			95		•		•	100	_				105	
GAA	AGA	AGT	TAC	TGG	AAA	TCT	CAA	GTC	AAC	CAT	GAT	ATT	TGC	AGG	953
											Asp				
	_		•	110	•				115		-			120	
ГСТ	GAG	CCG	GTC	TCT	GCT	GCG	GTG	TTG	AGG	AAA	CTC	CAG	GTG	GAA	998
											Leu				
				125					130	•				135	
rgg	CCC	TTC	CAG	GAG	CTG	TTG	CTG	CTC	CGA	GAG	CAG	TAC	CAA	AAG	1043
											Gln				
				140					145			-		150	
SAG	TTC	AAG	AAC	AGC	ACC	TAC	TCA	AGA	AGC	TCA	GTG	GAC	ATG	CTC	1088
											Val				
		-		155	-	-	_	_	160			-		165	
CAC	AGT	TTT	GÇC	AAG	TGC	TCG	GGG	TTA	GAC	CTG	ATC	TTT	GGT	CTA	1133
											Ile				
-				170		_	•		175					180	
									-					-	
AT	GCG	TTA	CTA	CGA	ACC	CCA	GAC	TTA	CGG	TGG	AAC	AGc	TCC	AAC	1178
											Agn				

		34	
	185	190	195
		GC TCT TCC AAG GGT ys Ser Ser Lys Gly	
Ala Gin Deu Deu	200	ys ser ser bys Gry	lyr Asn lle 210
	, =00	203	210
TCC TGG GAA CTG	GGC AAT GAG C	CC AAC AGT TTC TGG	AAG AAA GCT 1268
Ser Trp Glu Leu	Gly Asn Glu P	ro Asn Ser Phe Trp	Lys Lys Ala
	215	220	225
CAC ATT CTC ATC	CAM COO MING C	AC META CON CON CON	
		AG TTA GGA GAA GAC : ln Leu Gly Glu Asp 1	
	230	235	240
			240
TTG CAT AAA CTT	CTA CAA AGG TO	CA GCT TTC CAA AAT O	CA AAA CTC 1358
Leu His Lys Leu	Leu Gln Arg Se	er Ala Phe Gln Asn A	la Lys Leu
	245	250	255
TAT GGT CCT GAC	ATC CCT CAC C	CT CGA GGG AAG ACA G	mm
		o Arg Gly Lys Thr V	
-1, ··	260	265	270
CTG AGG AGT TTC	CTG AAG GCT GG	C GGA GAA GTG ATC G	AC TCT CTT 1448
		y Gly Glu Val Ile A	sp Ser Leu
	275	280	285
ACA TGG CAT CAC	ቸልጥ ጥልሮ ምፕር ል ል	T GGA CGC ATC GCT A	CC AAA GAA 1493
		n Gly Arg Ile Ala T	
	290	295	300
		G GAC ACT TTT ATT C	
		u Asp Thr Phe Ile L	
,	305	310	315
CAA AAA ATT CTG	AAG GTC ACT AA	A GAG ATC ACA CCT G	GC AAG AAG 1583
		s Glu Ile Thr Pro G	
;	320	325	330
		A GCT TAC GGT GGC GG	
	310 Thr Ser Sei 335	r Ala Tyr Gly Gly G: 340	
•	,,,,	340	345
TTG CTG TCC AAC A	ACC TTT GCA GCT	GGC TTT ATG TGG CT	G GAT AAA 1673
Leu Leu Ser Asn 7	Thr Phe Ala Ala	Gly Phe Met Trp Le	u Asp Lys
3	350	355	360
MING COO CING MOS C			
		: ATA GAA GTC GTG AT · Ile Glu Val Val Me	
	ia din Mec diy	370	375
_		5.0	373
GTG TTC TTC GGA G	CA GGC AAC TAC	CAC TTA GTG GAT GA	A AAC TTT 1763
Val Phe Phe Gly A	la Gly Asn Tyr	His Leu Val Asp Gl	u Asn Phe
3	80	385	390
CAC CCT TOTA CCT C	NT TNC TCC	TOT COM	0 111 0mg
		TCT CTT CTG TTC AA Ser Leu Leu Phe Ly	
	95	400	405
•			

GTA	GGI	, CC	C AG	G GT	G TT/	CTC	TC	A AG	A GT	G AA	A GG	c cc	A GA	C AGG	1853
Val	Gly	Pre	o Ar	g Va	l Lei	. Leu	Sez	Arg	y Va	l Lys	s Gl	y Pr	o As	p Arg	
				410	ס				415	5				420	
														C CCA	1898
Ser	Lys	Leı	ı Ar			Leu	Hie	су Су	Thi	: Asr	ı Va	l Ty	r Hi	s Pro	
				425	5				430)				435	
CCA	77 h m	-													
														CAT	1943
Arg	ıyı	GII	ı GII	440		Leu	Thr	Leu			. Let	ı Ası	ı Leı	ı His	
				440					445	,				450	
TAA	GTC	ACC	. 220	ב ראר	TTC	220	COL		com		mme			AAA	
				His											1988
			-,.	455	Deu	Lys	Val	FIU	460	PIO	ner	Pile	a Arc	465	
									400					405	
CCA	GTG	GAT	ACG	TAC	CTT	CTG	AAG	ССТ	TCG	GGG	CCG	CAT	· ccz	מידים ו	2033
				Tyr											2033
				470			•	-	475	3				480	
CTT	TCC	AAA	TCT	GTC	CAA	CTG	AAC	GGT	CAA	ATT	CTG	AAG	ATG	GTG	2078
				Val											
				485				_	490			-		495	
GAT	GAG	CAG	ACC	CTG	CCA	GCT	TTG	ACA	GAA	AAA	CCT	CTC	ccc	GCA	2123
				Leu											
				500					505					510	
				AGC											2168
Gly :	Ser	Ala	Leu	Ser	Leu	Pro .	Ala	Phe	Ser	Tyr	Gly	Phe	Phe	Val	
				515					520					525	
ATA A										TGA	AAA	TAA	AAG	GCA	2213
Ile /	Arg	Asn	Ala	Lys	Ile .	Ala	Ala	Сув	Ile						
				530					535						
TAC															2258
ACA A															2303
CTT															2348
TTC 1	ici (-IA	AGA	AGA A	ATA (TG C	CAG (GTG (GTG A	ACA (GTT	AAT	AGC	ACT	2393
GIG															2396
(2)	τ.	NPOE	MA TO	ON F	OD 0	PO T	D 110								
(2)		i)		EQUE											
	,	-,		EQUE: A)		NGTH		CISI	385						
				B)		NGIA PE:	•								
				C)		PE: RANDI	FDMF	cc.		leic	acı	.a			
				D)		POLO		JJ :	lin						
	(:	ki)		EQUEN				ON-			NO - 1	•			
CGGCC													CD	50	
AGGCA															
TTTAC														100 150	
ACCAT	CGAC	G CC	AGT	TGGC	CAC	CGAC	CCT	CGGT	TCCT	ירא כ		اراناسار. مردوع	A.C.	200	
CTCTC														250	
GATTT	GGCG	G CA	CCA	GACT	GAC	TTCC'	TTA	L'Trim	TGAT	יכר כ	ים ממי	PCC.	22	300	
CCCAC														350	

36

TTGCGGGTCT	GACCGGGTCT	CCGCTGACGT	GTTGA

385

		•				-
NFORMA	TION FO	R SEQ ID NO	0:47:			
(i)	SEQUENC	E CHARACTE	RISTI	CS:		
	(A)	LENGTH:		541		
	(B)	TYPE:		nuclei	c acid	
	(C)	STRANDEDNE	SSS:	double		
	(D)	TOPOLOGY:		linear		
xi)	SEQUENC	E DESCRIPT	ION:	SEQ ID	NO:47	
AC ATAT	CCTTCA	CTTATTTGCC	TCTTC	GTCAT	ATTGGAGGCA	50
CA TTTT	TAATAA	CCCTCAAAAT	AGTGC	ATGCA	AAGTGCTAAG	100
CACA	TGGTGC (CATTAACTGT	CACCA	CCTGC	AGTGGTCTAC	150
CA CCGC	ACTGGA :	TGTTAACACT	GAAGO	CCGTG	CCCCGCCCTC	200
T GGAT	CCAGCG :	TTGAAGCTTG	CCCCG	CCCTC	CCGAGGCTCT	250
A CTGG	AGCATG (CCCGCCCTC	CCGAG	GCTCT	GGAGCTTGCT	300
G CTCC	CTACCG (TGGGGTTTT	GCTTT	ATTCT	TATGAATGAC	350
C GCTT	TCGTCT (CAGGGGTACT	GTAAT	GCCTT	TTATTTTCAT	400
G CGAT	TTTGGC A	TTTCTTATG	ACAAA	AAACC	CATAGGAAAA	450
G CTTA	GTGAGC 1	TTCCTGCGGG	GAGAG	GTTTT	TCTGTTAGAG	500
T CTGC	CATCG A	CCATCTTCA	GGCCT	CGTGC	c	541
	xi) AC ATAT AC ATAT AC CACA ACCGC T GGAT ACTGG CC CC GCTT CC CCTT CC CCTT CC CCTT	(A) (B) (C) (D) Xi) SEQUENC AC ATATCCTTCA CA TTTTTAATAA CC CACATGGTGC CA CCGCACTGGA CT GGATCCAGCG CA CTGGAGCATG CG CTCCCTACCG CG CTCCCTACCG CG CGATTTTGGC CG CGATTTTGGC CG CGATTTTGGC CG CTTAGTGAGC CT	(A) LENGTH: (B) TYPE: (C) STRANDEDNE (D) TOPOLOGY: XI) SEQUENCE DESCRIPT. AC ATATCCTTCA CITATITGCC ATTITTAATAA CCCTCAAAAT C CACATGGTGC CATTAACTGT CA CTGGAGCATG CCCCCCCTC CA CTCCCTACCG CTGGAGGTTTT CC CTCCCTACCG CTGGGGTTTT CC CGCATTTGGC ATTICTTATG CC CTTAGTGAGC TTCCTGCGGG CCTTAGTGAGC TTCCTGCGGG CTTAGTGAGC TTCCTGCGGG CTTAGTGAGC TTCCTGCGGG CTTAGTGAGC TTCCTGCGGG CTTAGTGAGC TTCCTGCGGG	(A) LENGTH: (B) TYPE: (C) STRANDEDNESS: (D) TOPOLOGY: XI) SEQUENCE DESCRIPTION: AC ATATCCTTCA CITATITGCC TCTTC CA TITITAATAA CCCTCAAAAT AGTGC CA CCGCACTGGA TGTTAACACT GAAGC TG GGATCCAGCG TTGAAGCTTG CCCCG CA CTGGAGCATG CCCCGCCCTC CCGAG CA CTGGAGCATG CCCCGCCTC CCGAG CA CTGCATCCG CTGGGGTTTT GCTTT CC GCTTTCGTCT CAGGGGTACT GTAAT CC CGATTTTGGC ATTTCTTATG ACAAA CG CTTAGTGAGC TTCCTGCGGG GAGAG	(A) LENGTH: 541 (B) TYPE: nuclei (C) STRANDEDNESS: double (D) TOPOLOGY: linear XI) SEQUENCE DESCRIPTION: SEQ ID AC ATATCCTTCA CITATITECC TCTTGGTCAT CA CTTATAATAA CCCTCAAAAT AGTGCATGCA CC CACATGGTGC CATTAACTGT CACCACCTGC CA CTGGAGCATG TGTTAACACT GAAGCGCGTG CT GGATCCAGCG TTGAAGCTT CCCCGCCCTC CA CTGGAGCATG CCCCGCCCTC CA CTGGAGCATG CCCCGCCCTC CA CTGCAGCATG CCCCGCCCTC CA CTGGAGCATG CCCCGCCCTC CC CTCCCTACCG CTGGGGTTTT GCTTTATTCT CC CGCTTTCGTCT CAGGGGTACT GTAATGCCTT CC CGTTTTTGGC ATTTCTTATG ACAAAAAACC CC CTTAGTGAGC TTCCTGCGGG GAGAGGTTTT	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 541 (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/03542

A. CLASSIFICATION OF SUBJECT MATTER IPC(7) :C12N 15/56, 15/63, 1/21, 9/24, 15/11 US CL :536/23.1, 23.2; 435/200, 325, 252.3, 320.1; 424/94.61 According to International Patent Classification (IPC) or to both national classification and IPC							
B. FIELDS SEARCHED							
Minimum documentation searched (classification system follo	Minimum documentation searched (classification system followed by classification symbols)						
U.S. : 536/23.1, 23.2; 435/200, 325, 252.3, 320.1; 424/94.61							
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched							
Electronic data base consulted during the international search	(name of data base and, where practi	sable search terms used)					
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EAST, MEDLINE, BIOSIS, CAPLUS, SCISEARCH, EMBASE, JAPIO, PATOWEP, PATOSWO search terms: heparanase, gene or sequence							
C. DOCUMENTS CONSIDERED TO BE RELEVANT		·					
Category* Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.					
X US 5,362,641 A (FUKS et al.) 0 document.	8 November 1994, see ent	ire 21-25					
Y		1-20, 26-28					
X, P US 5,968,822 A (PECKER et al.) document.	19 October 1999, see ent	ire 1-28					
X WO 95/04158 A1 (THE UPJOHN (WO 95/04158 A1 (THE UPJOHN COMPANY) 09 Febuary 1995,						
Y see entire document.	see entire document.						
X, P WO 99/11798 A1 (INSIGHT STRATE) 11 March 1999, see entire document	TEGY & MARKETING LTI	D.) 1-28					
9							
Further documents are listed in the continuation of Box	C. See patent family anne	к.					
Special categories of cited documents: 'A' document defining the general state of the art which is not considered to be of particular relevance		e international filing date or priority application but cited to understand as the invention					
B* earlier document published on or after the international filing date "L* document which may throw doubts on priority claim(s) or which is	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step						
cited to establish the publication date of another citation or other special reason (as specified) O' document referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is						
pe document published prior to the international filing date but later than	combined with one or more other such documents, such combination being obvious to a person skilled in the art *A* document member of the same patent family						
the priority date claimed Oute of the actual completion of the international search	search report						
12 JUNE 2000	24 JUL 2000						
lame and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	Authoritod officer RICHARD HUTSON Authoritod Successed Jou						
acsimile No. (703) 305-3230	Telephone No. (703) 308-0196						

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/03542

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)							
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:							
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:							
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements t an extent that no meaningful international search can be carried out, specifically:	o such						
Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4. Claims Nos.:	(a).						
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)							
This International Searching Authority found multiple inventions in this international application, as follows:	0						
Please See Extra Sheet.							
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all claims.	scarchable						
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invit of any additional fee.	e payment						
3. As only some of the required additional search fees were timely paid by the applicant, this international search reports those claims for which fees were paid, specifically claims Nos.:	oort covers						
4. No required additional search fees were timely paid by the applicant. Consequently, this international search restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	n report is						
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.							
No protest accompanied the payment of additional search fees.							

Form PCT/ISA/210 (continuation of first sheet(1)) (July 1998)*

INTERNATIONAL SEARCH REPORT

International application No. PCT/US00/03542

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claim(s)1-7, 19, 20 and 28, drawn to a nucleic acid encoding a polypeptide having heparanase activity.

Group II, claim(s) 8-18, drawn to antisense oligonucleotides of a polynucleotide which encodes a polypeptide having heparanase activity.

Group III, claim(s) 21-25, drawn to polypeptide having heparanase activity.

Group IV, claim(s)26, drawn to a method of identifying a chromosome region harboring a heparanase gene.

Group V, claim(s) 27, drawn to a method of eliciting anti-heparanase antibodies in vivo.

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons: The listed inventions share a technical relationship of a polypeptide having heparanase activity, but this does not constitute a special technical feature because Fuks et al. (Fuks et al. US Patent No: 5,362,641) teach a polypeptide having heparanase activity.